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FILING DATE: August 26, 2002

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### TELEPHONE 510-923-3179 USE ONLY FOR FILING A PROVISIONAL APPLICATION FOR PATENT

Docket Number:

19195.001

This collection of information is required by 37 CFR 1.51. The information is used by the public to file (and by the PTO to process) a provisional application. Confidentiality is governed by 35 U.S.C. 122 and 37 CFR 1.14. This collection is estimated to take 8 hours to complete, including gathering, preparing, and submitting the complete provisional application to the PTO. Time will vary depending upon the individual case. Any comments on the amount of time y u require to complete this form and/or suggestions for reducing this burden, should be sent to the Chief Information Officer, U.S. Patent and Trademark Office, U.S. Department of Commerce, Washington, D.C. 20231. DO NOT SEND FEES OR COMPLETED FORMS TO THIS ADDRESS. SEND TO: Box Provisional Application, Assistant Commissioner for Patents, Washington, D.C. 20231.

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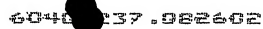
# PROVISIONAL APPLICATION COVER SHEET Additional Page

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Serial No. Filing Date To be assigned 08/26/2002		Examiner To be assigned	Group Art Unit To be assigned	
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#### CONSERVED AND SPECIFIC STREPTOCOCCAL GENOMES

#### FIELD OF THE INVENTION

The invention relates to polynucleotides which are conserved or specific to one or more species of Streptococcus, Streptococcus species serotypes, and/or serotype isolates. The conserved or specific genomic regions can be used to identify, screen and develop vaccines and other treatments for Streptococcal infections and can be used in diagnostic assays to diagnose and identify Streptococcal infections.

#### **BACKGROUND OF THE INVENTION**

The genus *Streptococcus* consists of Gram-positive, chain-forming, spherical bacterial cells. Three species of clinical interest are *S.pneumoniae* ("pneumococcus" or "S.pn."), *S.pyogenes* ('group A streptococcus' or 'GAS') and *S.agalactiae* ('group B streptococcus' or 'GBS'). Infections with these three pathogenic streptococci lead to conditions including pharyngitis, toxic shock syndrome and necrotizing fasciitis.

Once thought to infect only cows, GBS is now known to cause serious disease, bacteraemia and meningitis in immunocompromised individuals and neonates. There are two known types of neonatal infection. The first (early onset, usually within 5 days of birth) is manifested by bacteraemia and infection. It is generally contracted vertically as a baby passes through the birth canal. GBS is thought to colonize the vagina of about 25% of young women; approximately 1% of infants born via a vaginal birth to colonised mothers will become infected. Mortality resulting from these infections is between 50 - 70%. The second type of neonatal infection is a meningitis that occurs 10 to 60 days after birth. If pregnant women are vaccinated with type III capsule so that the infants are passively immunised, the incidence of the late onset meningitis is generally reduced, although not entirely eliminated.

The "B" in "GBS" refers to the Lancefield classification, which is based on the antigenicity of a carbohydrate which is soluble in dilute acid and called the C carbohydrate. Lancefield identified 13 types of C carbohydrate, designated A to O,



that could be serologically differentiated. The organisms that most commonly infect humans are found in groups A, B, D, and G. Within group B, strains can be divided into at least 9 serotypes (Ia, Ib, II, III, IV, V, VI, VII, and VIII) based on the structure of their polysaccharide capsule. Further categories based on, for example, the expression of certain proteins have also been developed.

GBS strains of polysaccharide capsule Type V were rarely isolated before the mid-1980's but now account for approximately one-third of clinical isolates in the US. Type V is the most common capsular serotype associated with invasive infection in nonpregnant adults, and the emergence of Type V strain over the past decade has been temporarily linked to an increase in GBS disease in this population.

Group A streptococcus is a frequent human pathogen, estimated to be present in between 5-15% of normal individuals without signs of disease. When host defences are compromised, or when the organism is able to exert its virulence, or when it is introduced into vulnerable tissues or hosts, however, an acute infection occurs. Diseases include puerperal fever, scarlet fever, erysipelas, pharyngitis, impetigo, necrotising fasciitis, myositis and streptococcal toxic shock syndrome.

Pneumococcus is the most common cause of acute respiratory infection and otitis media and is estimated to result in over 3 million deaths in children every year worldwide from pneumonia, bacteremia, or meningitis. Even more deaths occur among elderly people, among whom S. pn. is the leading cause of community-acquired pneumonia and meningitis. Since 1990, the number of penicillin-resistant strains has increased from 1 to 5% to 25 to 80% of isolates, and many strains are now resistant to commonly prescribed antibiotics such as penicillin, macrolides, and fluoroquinolones. See Tettelin, et al. (2001) Science 293, 248-506.

The complete genomic sequence of a virulent isolate of *S. pneumoniae* was published by Tettelin, et al. (2001) *Science* 293, 248-506 and is available at the TIGR website at http://www.tigr.com. The genomic sequence, the Tettelin article and its published supplemental material are incorporated herein by reference in their entirety.

The complete genomic sequence of an M1 strain of S. pyrogenes was published by Ferretti, et al. (2001) Proc. Natl. Acad. Sci. USA 98, 4658 – 4663 and is available at the TIGR website at http://www.tigr.com. The genomic sequence, the Ferretti article and its published supplemental materials are incorporated herein by reference in their entirety.

The complete genomic sequence of a serotype V strain of S. agalactiae (type V strain 2603 V/R) is published on the date of this filing, August 26, 2002 by Tettelin, et al. (2002) Proc. Natl. Acad. Sci. USA and/or is available on the same day at the TIGR website at http://www.tigr.com. Most of this sequence is also available in PCT International Patent Application Publication WO 02/34771. The genomic sequence, the Tettelin article and its published supplemental materials are incorporated herein by reference in their entirety.

Current treatments for *Streptococcal* infections include both antibiotics and prophylactic vaccination. Current vaccines, particularly with respect to GBS, suffer from poor immunogenicity, while the emergence of antibiotic resistant strains has lessened the effectiveness of currently used antibiotics. Accordingly, there is an increasing need for the development of new vaccines and antibiotics (as well as other small molecule bacterial inhibitors) to help prevent and treat Streptococcal infections.

Applicants have identified regions of the Streptococcal genomes which can be used to identify and develop new vaccines and treatments for Streptococcal infections. Specifically, Applicants have identified polynucleotides of the Streptococcal genome which are conserved or specific to Streptococcal species, species serotypes, and/or specific serotype isolates. These polynucleotides and their expressed polypeptides can be used to screen, develop and design new vaccines, antibiotics and other small molecule bacterial inhibitors. These polynucleotides and their expressed polypeptides can further be used to diagnose and identify Steptococcal infections.

#### SUMMARY OF THE INVENTION

The invention relates to polynucleotides which are conserved or specific to one or more species of Streptococcus, Streptococcus species serotypes, and/or serotype isolates. In particular, the invention relates to polynucleotides from Streptococcus which are conserved or specific to one or more of the species of S. pneumoniae ("pneumococcus" or "S. pn."), S. pyogenes ("group A streptococcus" or "GAS"), and S. agalactiae ("group B streptococcus" or "GBS"). The invention further relates to polynucleotides which are conserved or specific to one or more Streptococcal species serotypes, such as GBS serotypes Ia, Ib, II, III, IV, V, VI, VII, and VIII. The invention still further relates to polynucleotides which are conserved or specific to one or more clinical isolates of a Streptococcus species.

The invention is based on the identification of the following Subsets of genes. Genes falling within each subset are described with respect to referenced tables, lists, and/or figures (in particular the CGH map depicted in Figure 1).

The following Subsets related to the GBS genome:

GBS Subset 1: 1060 GBS genes which have homologs with GAS and with pneumococcus (Table 8);

GBS Subset 2: 225 GBS genes which have homologues with GAS, but not with pneumococcus (Table 10);

GBS Subset 3: 176 GBS genes which have homologues with pneumococcus but not with GAS (Table 9);

GBS Subset 4: 683 GBS genes which do not have homologues with GAS or pneumococcus (specific to GBS vs GAS and pneumococcus) (Table 11).

The invention is based on the identification of the following subsets of genes within the GAS genome:

GAS Subset 1: 1006 GAS genes which have homologues with GBS and with pneumococcus (Table 33);

GAS Subset 2: 212 GAS genes which have homologues with GBS but do not have homologues with pneumococcus (Table 34);

GAS Subset 3: 62 GAS genes which have homologues with pneumococcus but do not have homologues with GBS (Table 35);

GAS Subset 4: 416 GAS genes which do not have homologues with either GBS or pneumococcus. This Subset can be determined by subtracting the above subsets from the published genome.

The invention is based on the identification of the following subsets of genes within the pneumococcus genome:

Spn Subset 1: 1034 Spn genes which have homologues with GBS and GAS (Table 36);

Spn Subset 2: 195 Spn genes which have homologues with GBS but do not have homologues with GAS (Table 37);

Spn Subset 3: 74 Spn genes which have homologues with GAS but do not have homologues with GBS (Table 38);

Spn Subset 4: 836 Spn genes which do not have homologues with either GBS or pneumococcus. This Subset can be determined by substracting the above Subsets from the published genome.

The invention further provides polynucleotides which are conserved or specific to Streptococcus based on a comparison with a wide range of published bacterial genomes. The following additional Subsets are provided:

GBS Subset 1(a): Of the 1060 GBS genes which have homologues in both GAS and pneumococcus, 12 of those GBS genes do not have homologues with any of the other published bacterial genomes at the time of the invention (i.e., GBS Subset 1(a) is specific to Streptococcus vs non Streptococcus published genomes). (The 12 GBS ORF's are listed in Table 3).

GBS Subset 2(a): This Subset comprises GBS genes which have homologues with GAS, but not with pneumococcus or any other published bacterial genomes at the time of the invention.

GBS Subset 3(a): This Subset comprises GBS genes which have homologues with pneumococcus, but not with GAS or any other published bacterial genomes at the time of the invention.

GBS Subset 4(a): Of the 683 GBS genes which do not have homologues in either GAS or pnuemococcus, 315 of these GBS genes also do not have homologues with any of the other published bacterial genomes. These include six proteins predicted to be anchored on the cell wall (SAG0677, SAG0771, SAG1052, SAG1331, SAG1473, and SAG1168), three of the capsule-related genes (SAG1163, SAG1167, and SAG1168), six transcriptional regulators, and four genes of the cyl operon (SAG0663 – SAG0673) essential for GBS hemolytic activity and production of pigment. See Pritzlaff et al. (2001) *Mol. Microbiol.*, 39, 236 – 247. The rest of the 315 proteins include 240 hypothetical proteins with no similarity to other proteins in databases.

Many of the 315 genes specific to *S. agalactiae* are located in regions likely to constitute mobile genetic elements. Two of these regions resemble prophages (SAG0545-SAG0610 and SAG1835-SAG1885) displaying a mosaic structure with segments most similar to different bacteriophages, a pattern that suggests frequent recombination events. PblA and PblB are adhesins from a *S. mitis* prophage where they contribute to endocarditis by binding to human platelets (See Bensing, et al. (2001) *Infect. Immun.* 69, 6186 – 6192; Bensing, et al (2001) *Infect. Immun.* 69, 1373 – 1380. Their orthologs in *S. agalactiae* are located on separate prophages and display a different protein structure. Another region (SAG1247-SAG1299) encodes a putative conjugative transposon that carries genes for cadmium efflux and mercury resistance.

GAS Subset 1(a): This Subset comprises GAS genes which have homologues with GBS and with pneumococcus, but do not have homologues with any of the other published bacterial genomes at the time of the invention.

GAS Subset 2(a): This Subset comprises GAS genes which have homologues with GBS but do not have homologues with pneumococcus or any of the other published bacterial genomes at the time of the invention:

GAS Subset 3(a): This Subset comprises GAS genes which have homologues with pneumococcus but do not have homologues with GBS or any of the other published bacterial genomes at the time of the invention.

GAS Subset 4(a): This Subset comprises GAS genes which do not have homologues with either GBS or pneumococcus or with any of the other published bacterial genomes at the time of the invention.

Spn Subset 1(a): This Subset comprises Spn genes which have homologues with GBS and GAS but which do not have homologues with any of the other published bacterial genomes at the time of the invention;

Spn Subset 2(a): This Subset comprises Spn genes which have homologues with GBS but do not have homologues with GAS or with any of the other published bacterial genomes at the time of the invention;

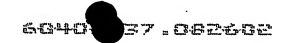
Spn Subset 3(a): This Subset comprises Spn genes which have homologues with GAS but do not have homologues with GBS or with any of the other published bacterial genomes at the time of the invention;

Spn Subset 4(a): This Subset comprises Spn genes which do not have homologues with either GBS or pneumococcus or with any of the other published bacterial genomes at the time of the invention.

The invention also provides polynucleotides which are conserved or specific to GBS serotypes and/or clinical isolates. Applicants have sequenced 19 GBS genes from a variety of GBS serotypes in 11 different clinical isolates. The sequences of these genes are set forth in Tables 13 - 31. The following additional subsets are provided:

GBS Subset 1(b): of the 1060 GBS genes which have homologues with GAS and with pneumococcus, 47 of these GBS genes vary among the 11 clinical isolates.

1013 of these GBS genes are conserved across the 11 clinical isolates. This list can



be determined by comparing the genes listed in Table 8 with the Comparative Genome Hybridization in Figure 1.

GBS Subset 2(b): of the 225 GBS genes which have homologues with GAS, but not pneumococcus, 44 of these GBS genes vary among the 11 clinical isolates.

181 of these GBS genes are conserved across the 11 clinical isolates. This list can be determined by comparing the genes listed in Table 10 with the Comparative Genome Hybridization in Figure 1.

GBS Subset 3(b): of the 176 GBS genes which have homologues with pneumococcus, 44 of these GBS genes vary among 11 clinical isolates. 132 of these GBS genes are conserved across the 11 clinical isolates. This list can be determined by comparing the genes listed in Table 9 with the Comparative Genome Hybridization in Figure 1.

GBS Subset 4(b): of the 683 GBS genes which do not have homologues with GAS or pneumococcus, 260 GBS genes vary among the 11 clinical isolates. 423 of these GBS genes are conserved across the 11 clinical isolates. This list can be determined by comparing the genes listed in Table 11 with the Comparative Genome Hybridization in Figure 1.

The invention further provides polynucleotides which are likely recent genomic duplications in GBS. These duplications include glycosyl transferases, sortases, proteins anchored on the cell wall, & lactam resistance factors, and many hypothetic proteins. The GBS genes are listed in Table 4 (GBS Subset 5).

The invention is also based on the identification of a cluster of 13 adjacent genes (SAG1410 – SAG1424) which is believed to encode enzymes required for synthesis of the group B carbohydrate, a coplex multiantennary structure of rhamnose, glucitol phosphate, N-acetylglucosamine, and galactose. (GBS Subset 6). Predicted proteins encoded within this cluster include seven putative glycoslytransferases, four of which are similar to rhamnosyltransferases in other streptococcal species; a putative dTDP-L-rhamnose synthase; and proteins involved in glucitol synthesis. All nine regonized GBS capsular polysaccharide types contain sialic acid residues as part

of their repeating unit structure, a feature that contributes to virulence by inhibiting activation of the alternative complement pathway. See Edwards et al. (1982) J. Immunol. 128, 1278 - 1283.

The type V capsular polysaccharide gene cluster consists of 18 genes. (GBS Subset 6(a)). A region of glycosyltransferases and related proteins (SAG1162 – SAG1170) that direct the synthesis of the type V polysaccharide repeat unit is flanked on either side by genes that are conserved in all known GBS capsule serotypes. Downstream of this region are genes that encode enzynmes for the biosynthesis and activation of sialic acid (SAG1158 – SAG1161). Upstream of the serotype specific region are genes (SAG1171 – SAG1175) found not only in all nine GBS capsular serotypes but also in a variety of other polysaccharide-producing streptococci.

The invention provides for methods of screening a Streptococcal genome for a conserved or a specific genomic sequence using one or more of the subsets of the invention.

The invention further provides for an immunogenic composition comprising a polypeptide expressed by one or more of the polynucleotides in one or more of the subsets of the invention, and methods for designing an immunogenic composition by selecting one or more polypeptides expressed by one or more of the polynucleotides in one or more of the subsets of the invention.

The invention further provides for methods of screening compounds for activity against a Streptococcal bacteria, which method comprises contacting the compounds with a polypeptide expressed by the polynucleotide from one of the subsets of the invention.

The invention further provides for compositions comprising one or more of the polynucleotides, and fragments and derivatives thereof, selected from the group consisting of the sequences set forth in Tables 13-31.

The invention further provides for compositions comprising polypeptides and fragments and derivatives thereof encoded by the polynucleotides set forth in Tables 13-31.

#### BRIEF DESCRIPTION OF THE TABLES AND DRAWINGS

<u>Table 1</u> comprises a complete list of GBS predicted genes, listed by SAGxxxx ORF number. This table also includes the predicted amino acid size of the predicted expressed protein and the predicted function, if known.

<u>Table 2</u> comprises a list of predicted and experimentally characterized surface and secreted proteins from GBS.

<u>Table 3</u> lists genes which were shared among GBS, GAS and pneumococcus, but which were not found in any of the other completely sequenced genomes.

Table 4 depicts probable recently duplicated genes within GBS.

<u>Table 5</u> lists the 19 GBS strains used for comparative genome hybridisations and phylogenetic analysis.

<u>Table 6</u> lists clusters of genes derived from phylogenetic profiling of GBS strains based on comparative genome hybridisations.

<u>Table 7</u> lists genes and strains used for phylogenetic analyses of the 19 GBS strains.

<u>Table 8</u> lists the 1060 GBS ORF's which are conserved across GBS, GAS and pneumococcus.

<u>Table 9</u> lists the 176 GBS ORF's which are conserved across GBS and pneumococcus.

Table 10 lists the 225 GBS ORF's which are conserved across GBS and GAS.

<u>Table 11</u> lists 683 GBS ORF's which are not shared with either GBS or pneumococcus.

<u>Table 12</u> lists 315 GBS ORF's which are not shared with any published genomic sequence.

Table 13 lists the sequences of the 11 strains relating to GBS ORF SAG0466.

Table 14 lists the sequences of the 11 strains relating to GBS ORF SAG0471.

Table 15 lists the sequences of the 11 strains relating to GBS ORF SAG0492.

Table 16 lists the sequences of the 11 strains relating to GBS ORF SAG0767.

Table 17 lists the sequences of the 11 strains relating to GBS ORF SAG1086.

Table 18 lists the sequences of the 11 strains relating to GBS ORF SAG1600.

Table 19 lists the sequences of the 11 strains relating to GBS ORF SAG1680.

Table 20 lists the sequences of the 11 strains relating to GBS ORF SAG1723.

Table 21 lists the sequences of the 11 strains relating to GBS ORF SAG0079.

Table 22 lists the sequences of the 11 strains relating to GBS ORF SAG0093.

Table 23 lists the sequences of the 11 strains relating to GBS ORF SAG0163.

Table 24 lists the sequences of the 11 strains relating to GBS ORF SAG0290.

Table 25 lists the sequences of the 11 strains relating to GBS ORF SAG0368.

Table 26 lists the sequences of the 11 strains relating to GBS ORF SAG0503.

Table 27 lists the sequences of the 11 strains relating to GBS ORF SAG1473.

Table 28 lists the sequences of the 11 strains relating to GBS ORF SAG1552.

Table 29 lists the sequences of the 11 strains relating to GBS ORF SAG1641.

Table 30 lists the sequences of the 11 strains relating to GBS ORF SAG2147.

Table 31 lists the sequences of the 11 strains relating to GBS ORF SAG2148.

<u>Table 32</u> provides a conversion table for the ORFxxxx reference numbers to the SAGxxxx reference numbers, which are available at the TIGR website on the day of the filing of this application.

<u>Table 33</u> lists the 1006 GAS ORF's which are shared with GBS and Spn. The genes corresponding to these ORFs were published in GenBank; the numbers for the GAS ORF refer directly to their GenBank entries.

Table 34 lists the 212 GAS ORF's which are shared with GBS but which do not have homologues with pneumococcus. The genes corresponding to these ORFs were published in GenBank; the numbers for the GAS ORF refer directly to their GenBank entries.

Table 35 lists the 62 GAS ORF's which have homologues with pneumococcus but which do not have homologues with GBS. The genes corresponding to these ORFs were published in GenBank; the numbers for the GAS ORF refer directly to their GenBank entries.

Table 36 lists the 1034 Spn ORF's which share homologues with GBS and GAS. These ORF's were published in GenBank. The numbers for Spn correspond to the entry for AE005672.

Table 37 lists the 195 Spn ORF's which share homologues with GBS but do not share homologues with GAS. These ORF's were published in GenBank. The numbers for Spn correspond to the entry for AE005672.

Table 38 lists the 74 Spn ORF's which share homologues with GAS but do not share homologues with GBS. These ORF's were published in GenBank. The numbers for Spn correspond to the entry for AE005672.

<u>Figure 1</u> is a circular representation of the GBS genome and comparative hybridisations using microarrays.

<u>Figure 2</u> is a schematic representation of in silico comparisons between streptococci.

Figure 3 depicts a phylogenetic tree of GBS strains based on PCR sequences.

Figure 4 depicts a linear representation of the GBS genome.

<u>Figure 5</u> demonstrates phylogenetic profiling of GBS strains based on comparative genome hybridisations.

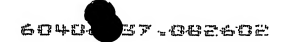
#### BRIEF DESCRIPTION OF THE SEQUENCE ID NOS.

The following SEQ ID NOS are used in the application and figures.

SEQ ID NOS. 1301 – 1316 represent the polynucleotide sequences corresponding to the SAG0466 ORF (thiolase) in the GBS strains indicated for each sequence, including where indicated reverse complements.

SEQ ID NOS. 1401 – 1417 represent the polynucleotide sequences corresponding to the SAG0471 ORF (glucokinase) in the GBS strains indicated for each sequence, including where indicated reverse complements.

SEQ ID NOS. 1501 – 1511-represent the polynucleotide sequences corresponding to the SAG0492 ORF (amino acid ABC transporter, ATP-binding



protein) in the GBS strains indicated for each sequence, including where indicated reverse complements.

SEQ ID NOS. 1601 – 1617 represent the polynucleotide sequences corresponding to the SAG0767 ORF (D-alanine - D-alanine ligase) in the GBS strains indicated for each sequence, including where indicated reverse complements.

SEQ ID NOS. 1701 – 1711 represent the polynucleotide sequences corresponding to the SAG1086 ORF (xanthine phosphoribosyltransferase) in the GBS strains indicated for each sequence, including where indicated reverse complements.

SEQ ID NOS. 1801 - 1814 represent the polynucleotide sequences corresponding to the SAG1600 ORF (glutamate racemase) in the GBS strains indicated for each sequence, including where indicated reverse complements.

SEQ ID NOS. 1901 – 1914 represent the polynucleotide sequences corresponding to the SAG1680 ORF (shikimate 5-dehydrogenase) in the GBS strains indicated for each sequence, including where indicated reverse complements.

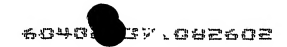
SEQ ID NOS. 2001 – 2010 represent the polynucleotide sequences corresponding to the SAG1723 ORF (signal peptidase I) in the GBS strains indicated for each sequence, including where indicated reverse complements.

SEQ ID NOS. 2101 – 2112 represent the polynucleotide sequences corresponding to the SAG0079 ORF (adenylate kinase) in the GBS strains indicated for each sequence, including where indicated reverse complements.

SEQ ID NOS. 2201 – 2211 represent the polynucleotide sequences corresponding to the SAG0093 ORF (D-alanyl-D-alanine carboxypeptidase family protein) in the GBS strains indicated for each sequence, including where indicated reverse complements.

SEQ ID NOS. 2301 – 2311 represent the polynucleotide sequences corresponding to the SAG0163 ORF (competence protein Cg1A) in the GBS strains indicated for each sequence, including where indicated reverse complements.

SEQ ID NOS. 2401 – 2410 represent the polynucleotide sequences corresponding to the SAG0290 ORF (ABC transporter, substrate-binding protein) in



the GBS strains indicated for each sequence, including where indicated reverse complements.

SEQ ID NOS. 2501 – 2511 represent the polynucleotide sequences corresponding to the SAG0368 ORF (protein of unknown function) in the GBS strains indicated for each sequence, including where indicated reverse complements.

SEQ ID NOS. 2601 – 2609 represent the polynucleotide sequences corresponding to the SAG0503 ORF (lipase/acylhydrolase) in the GBS strains indicated for each sequence, including where indicated reverse complements.

SEQ ID NOS. 2701 – 2711 represent the polynucleotide sequences corresponding to the SAG1473 ORF (cell wall surface anchor family protein) in the GBS strains indicated for each sequence, including where indicated reverse complements.

SEQ ID NOS. 2801 – 2811 represent the polynucleotide sequences corresponding to the SAG1552 ORF (conserved hypothetical protein) in the GBS strains indicated for each sequence, including where indicated reverse complements.

SEQ ID NOS. 2901 – 2911 represent the polynucleotide sequences corresponding to the SAG1641 ORF (YaeC family protein) in the GBS strains indicated for each sequence, including where indicated reverse complements.

SEQ ID NOS. 3001 – 3010 represent the polynucleotide sequences corresponding to the SAG2147 ORF (protein of unknown function / lipoprotein, putative) in the GBS strains indicated for each sequence, including where indicated reverse complements.

SEQ ID NOS. 3101 – 3111 represent the polynucleotide sequences corresponding to the SAG2148 ORF (LysM domain protein) in the GBS strains indicated for each sequence, including where indicated reverse complements.

#### DETAILED DESCRIPTION OF THE INVENTION

The invention relates to polynucleotides which are conserved or specific to one or more species of Streptococcus, Streptococcus species serotypes, and/or



serotype isolates. In particular, the invention relates to polynucleotides from Streptococcus which are conserved or specific to one or more of the species of S. pneumoniae ("pneumococcus" or "S. pn."), S. pyogenes ("group A streptococcus" or "GAS"), and S. agalactiae ("group B streptococcus" or "GBS"). The invention further relates to polynucleotides which are conserved or specific to one or more Streptococcal species serotypes, such as GBS serotypes Ia, Ib, II, III, IV, V, VI, VII, and VIII. The invention still further relates to polynucleotides which are conserved or specific to one or more clinical isolates of a Streptococcus species.

In order to facilitate an understanding of the invention, selected terms used in the application will be discussed below.

As used herein, the phrase "species of Streptococcus" generally refers to species of the Streptoccus family, including S.pneumoniae ("pneumococcus" or "S.pn."), S.pyogenes ('group A streptococcus' or 'GAS') and S.agalactiae ('group B streptococcus' or 'GBS').

As used herein, the phrase "Streptococcus species serotypes" generally refers to subdivisions based on a distinguishing characteristic within a specific Streptococcus species. The distinguishing characteristic can be identified by any of a wide range of diagnostic tools. For instance, GBS is generally recognized as comprising at least nine subdividing serotypes based on the structure of their polysaccharide capsule.

As used herein, the phrases "<u>serotype isolates</u>" or "<u>clinical isolates</u>" generally refer to specific isolated bacterial strains of a specific Streptococcal species and serotype.

As used herein in reference to bacterial genomes, the phrases "conserved" or "shared" generally refer to genomic sequences which have homologues in the two or more genomes in the reference. Homologous sequences preferably have greater than 50% identity (e.g., 60%, 70%, 80%, 90%, 95%, 99% or more).

As used herein in reference to bacterial genomes, the phrases "specific to" or "not shared" generally refer to genomic sequences which do not have homologues in

the two or more genomes in the reference. Sequences which are not homologues preferably have less than 50% identity (e.g., 40%, 35%, 30%, 25%, 20%, 15%, or less).

Identity between nucleotide sequences can be determined using software programs known in the art, for example those described in section 7.7.18 of *Current Protocols in Molecular Biology* (F.M. Ausubel *et al.*, eds., 1987) Supplement 30. A preferred alignment program is GCG Gap (Genetics Computer Group, Wisconsin, Suite Version 10.1), preferably using default parameters, which are as follows: open gap = 3; extend gap = 1.

Sequences within a Subset of the invention include sequences which hybridize to the listed genes. Hybridization reactions can be performed under conditions of different "stringency". Conditions that increase stringency of a hybridization reaction of widely known and published in the art [e.g. page 7.52 of Sambrook et al. (1989) Molecular Cloning: A Laboratory Manual. NY, Cold Spring Harbor Laboratory]. Examples of relevant conditions include (in order of increasing stringency): incubation temperatures of 25°C, 37°C, 50°C, 55°C and 68°C; buffer concentrations of 10 x SSC, 6 x SSC, 1 x SSC, 0.1 x SSC (where SSC is 0.15 M NaCl and 15 mM citrate buffer) and their equivalents using other buffer systems; formamide concentrations of 0%, 25%, 50%, and 75%; incubation times from 5 minutes to 24 hours; 1, 2, or more washing steps; wash incubation times of 1, 2, or 15 minutes; and wash solutions of 6 x SSC, 1 x SSC, 0.1 x SSC, or de-ionized water. Hybridization techniques and their optimization are well known in the art [e.g. see Sambrook et al.; RNA Methodologies (Farrell, 1998) (Academic Press; ISBN 0-12-249695-7); Current Protocols in Molecular Biology (F.M. Ausubel et al., eds., 1987) Supplement 30; Short protocols in molecular biology (4th edition, 1999) Ausubel et al. eds. ISBN 0-471-32938-X; US patent 5,707,829 etc.].

Identity between polypeptide sequences can be determined using software programs known in the art, for example those described in section 7.7.18 of *Current Protocols in Molecular Biology* (F.M. Ausubel *et al.*, eds., 1987) Supplement 30. A

preferred alignment is determined by the Smith-Waterman homology search algorithm [Smith & Waterman (1981) Adv. Appl. Math. 2: 482-489.] using an affine gap search with a gap open penalty of 12 and a gap extension penalty of 2, BLOSUM matrix 62.

Typically, 50% identity or more between two proteins may be considered to be an indication of functional equivalence. References to a percentage sequence identity between two amino acid sequences means that, when aligned, that percentage of amino acids are the same in comparing the two sequences.

The terms "polypeptide", "protein" and "amino acid sequence" as used herein generally refer to a polymer of amino acid residues and are not limited to a minimum length of the product. Thus, peptides, oligopeptides, dimers, mulimers, and the like, are included within the definition. Both full-length proteins and fragments thereof are encompassed by the definition. Minimum fragments of polypeptides useful in the invention can be at least 3, 4, 5, 6, 7, 8, 9, 10, 11, 12, 13, 14, or even 15 amino acids. Typically, polypeptides useful in this invention can have a maximum length suitable for the intended application. Generally, the maximum length is not critical and can easily be selected by one skilled in the art.

Reference to polypeptides and the like also includes derivatives of the amino acid sequences of the invention. Such derivatives can include postexpression modifications of the polypeptide, for example, glycosylation, acetylation, phosphorylation, and the like. Amino acid derivatives can also include modifications to the native sequence, such as deletions, additions and substitutions (generally conservative in nature), so long as the protein maintains the desired activity. These modifications may be deliberate, as through site-directed mutagenesis, or may be accidental, such as through mutations of hosts which produce the proteins or errors due to PCR amplification. Furthermore, modifications may be made that have one or more of the following effects: reducing toxicity; facilitating cell processing (e.g., secretion, antigen presentation, etc.); and facilitating presentation to B-cells and/or T-cells.

"Fragment" or "Portion" as used herein refers to a polypeptide consisting of only a part of the intact full-length polypeptide sequence and structure as found in nature. For instance, a fragment can include a C-terminal deletion and/or an N-terminal deletion of a protein.

A "<u>recombinant</u>" protein is a protein which has been prepared by recombinant DNA techniques as described herein. In general, the gene of interest is cloned and then expressed in transformed organisms, as described further below. The host organism expressed the foreign gene to produce the protein under expression conditions.

The term "polynucleotide", as known in the art, generally refers to a nucleic acid molecule. A "polynucleotide" can include both double- and single-stranded sequences and refers to, but is not limited to, cDNA from viral, prokaryotic or eukaryotic MRNA, genomic RNA and DNA sequences from viral (e.g. RNA and DNA viruses and retroviruses) or prokaryotic DNA, and especially synthetic DNA sequences. The term also captures sequences that include any of the known base analogs of DNA and RNA, and includes modifications such as deletions, additions and substitutions (generally conservative in nature), to the native sequence, so long as the nucleic acid molecule encodes a therapeutic or antigenic protein. These modifications may be deliberate, as through site-directed mutagenesis, or may be accidental, such as through mutations of hosts that produce the antigens. Modifications of polynucleotides may have any number of effects including, for example, facilitating expression of the polypeptide product in a host cell. The term "polynucleotide" further includes DNA, RNA, DNA/RNA hybrids, DNA and RNA analogues such as those containing modified backbones (with modifications in the sugar and/or phosphates e.g. phosphorothioates, phosphoramidites etc.), and also peptide nucleic acids (PNA) and any other polymer comprising purine and pyrimidine bases or other natural, chemically or biochemically modified, non-natural, or derivatized nucleotide bases etc. Nucleic acid according to the invention can be prepared in many ways (e.g. by chemical synthesis, from genomic or cDNA libraries,

from the organism itself etc.) and can take various forms (e.g. single stranded, double stranded, vectors, probes etc.).

A polynucleotide can encode a biologically active (e.g., immunogenic or therapeutic) protein or polypeptide. Depending on the nature of the polypeptide encoded by the polynucleotide, a polynucleotide can include as little as 10 nucleotides, e.g., where the polynucleotide encodes an antigen.

By "<u>isolated</u>" is meant, when referring to a polynucleotide or a polypeptide, that the indicated molecule is separate and discrete from the whole organism with which the molecule is found in nature or, when the polynucleotide or polypeptide is not found in nature, is sufficiently free of other biological macromolecules so that the polynucleotide or polypeptide can be used for its intended purpose.

"Antibody" as known in the art includes one or more biological moieties that, through chemical or physical means, can bind to or associate with an epitope of a polypeptide of interest. The antibodies of the invention specifically bind to infectious prion conformations. The term "antibody" includes antibodies obtained from both polyclonal and monoclonal preparations, as well as the following: hybrid (chimeric) antibody molecules (see, for example, Winter et al. (1991) Nature 349: 293-299; and U.S. Patent No. 4,816,567; F(ab')<sub>2</sub> and F(ab) fragments; F<sub>v</sub> molecules (non-covalent heterodimers, see, for example, Inbar et al. (1972) Proc Natl Acad Sci USA 69:2659-2662; and Ehrlich et al. (1980) Biochem 19:4091-4096); single-chain Fv molecules (sFv) (see, for example, Huston et al. (1988) *Proc Natl Acad Sci USA* <u>85</u>:5897-5883); dimeric and trimeric antibody fragment constructs; minibodies (see, e.g., Pack et al. (1992) Biochem 31:1579-1584; Cumber et al. (1992) J Immunology 149B: 120-126); humanized antibody molecules (see, for example, Riechmann et al. (1988) Nature 332:323-327; Verhoeyan et al. (1988) Science 239:1534-1536; and U.K. Patent Publication No. GB 2,276,169, published 21 September 1994); and, any functional fragments obtained from such molecules, wherein such fragments retain immunological binding properties of the parent antibody molecule. The term

"antibody" further includes antibodies obtained through non-conventional processes, such as phage display.

As used herein, the term "monoclonal antibody" refers to an antibody composition having a homogeneous antibody population. The term is not limited regarding the species or source of the antibody, nor is it intended to be limited by the manner in which it is made. Thus, the term encompasses antibodies obtained from murine hybridomas, as well as human monoclonal antibodies obtained using human rather than murine hybridomas. See, e.g., Cote, et al. Monoclonal Antibodies and Cancer Therapy, Alan R. Liss, 1985, p 77.

An "immunogenic composition" as used herein refers to a composition that comprises an antigenic molecule where administration of the composition to a subject results in the development in the subject of a humoral and/or a cellular immune response to the antigenic molecule of interest. The immunogenicity of the composition may be facilitated by the use of an adjuvant. The immunogenic composition can be introduced directly into a recipient subject, such as by injection, inhalation, oral, intranasal or any other parenteral or mucosal (e.g., intra-rectally or intra-vaginally) route of administration.

The practice of the present invention will employ, unless otherwise indicated, conventional methods of chemistry, biochemistry, molecular biology, immunology and pharmacology, within the skill of the art. Such techniques are explained fully in the literature. See, e.g., Remington's Pharmaceutical Sciences, 18th Edition (Easton, Pennsylvania: Mack Publishing Company, 1990); Methods In Enzymology (S. Colowick and N. Kaplan, eds., Academic Press, Inc.); and Handbook of Experimental Immunology, Vols. I-IV (D.M. Weir and C.C. Blackwell, eds., 1986, Blackwell Scientific Publications); Sambrook, et al., Molecular Cloning: A Laboratory Manual (2nd Edition, 1989); Handbook of Surface and Colloidal Chemistry (Birdi, K.S. ed., CRC Press, 1997); Short Protocols in Molecular Biology, 4th ed. (Ausubel et al. eds., 1999, John Wiley & Sons); Molecular Biology Techniques: An Intensive Laboratory Course, (Ream et al., eds., 1998, Academic Press); PCR (Introduction to

Biotechniques Series), 2nd ed. (Newton & Graham eds., 1997, Springer Verlag); Peters and Dalrymple, Fields Virology (2d ed), Fields et al. (eds.), B.N. Raven Press, New York, NY.

It is understood that the antibodies and methods of this invention are not limited to particular formulations or process parameters as such may, of course, vary. It is also to be understood that the terminology used herein is for the purpose of describing particular embodiments of the invention only, and is not intended to be limiting.

All publications, patents and patent applications cited herein are hereby incorporated by reference in their entirety.

#### Vaccines and Immunisation

The invention provides an immunogenic composition comprising a polypeptide, or a fragment or derivative thereof, which is encoded by a polynucleotide sequence which is conserved across one or more species of Streptococcus.

The polynucleotide is preferably conserved across one or more species of Streptococcus selected from the group consisting of GBS, GAS and pneumococcus. In one embodiment, the polynucleotide is a GBS polynucleotide which is homologous with at least one gene from both GAS and pneumococcus. Preferably, the GBS polynucleotide is selected from GBS Subset 1, which includes 1060 GBS genes which have homologues with both GAS and pneumococcus (Table 8).

In another embodiment, the polynucleotide is a GAS polynucleotide which is homologous with at least one gene from both GBS and pneumococcus. Preferably, the GAS polynucleotide is selected from GAS Subset 1, which includes 1006 GAS genes which have homologues with both GBS and pneumococcus.

In another embodiment, the polypeptide is a pneumococcal polynucleotide which is homologous with at least one gene both GAS and GBS. Preferably, the pneumococcus polynucleotide is selected from Spn Subset 1, which includes 1034 pneumococcal genes which have homologous with both GBS and GAS.

In another embodiment, the polynucleotide is a GBS polynucleotide which is homologous with at least one gene from GAS. Preferably, the polynucleotide is selected from one of the genes listed GBS Subset 2, which includes 225 GBS genes which have homologues with GAS, but not with pneumococcus.

In another embodiment, the polynucleotide is a GBS polynucleotide which is homologous with at least one gene from pneumococcus. Preferably, the polynucleotide is selected from GBS Subset 3, which includes 176 GBS genes which have homologues with pneumococcus.

In another embodiment, the polynucleotide is a GAS polynucleotide which is homologous with at least one gene from GBS. Preferably, the polynucleotide is selected from GAS Subset 2, which includes 212 GAS genes which have a homologue with GBS.

In another embodiment, the polynucleotide is a GAS polynucleotide which is homologous with at least one gene from pneumoccus. Preferably, the polynucleotide is selected from GAS Subset 3, which includes 62 GAS genes which have a homologue with pneumococcus.

In another embodiment, the polynucleotide is a pneumococcus polynucleotide which is homologous with at least one gene from GBS. Preferably, the polynucleotide is selected from Spn Subset 2, which includes 195 Spn genes which have a homologue with GBS.

In another embodiment, the polynucleotide is a pneumococcus polynucleotide which is homologous with at least one gene from GAS. Preferably, the polynucleotide is selected from Spn Subset 3, which includes 74 Spn genes which have a homologue with GAS.

The invention further provides an immunogenic composition comprising a polypeptide, or a fragment or derivative thereof, which is encoded by a polynucleotide sequence which is specific to one or more species of Streptococcus.

The invention further provides an immunogenic composition comprising a polypeptide, or a fragment or derivative thereof, which is encoded by a polynucleotide

which is specific to GBS, GAS and pneumococcus. In one embodiment, the polynucleotide is a GBS polynucleotide which is homologous to at least one gene from both GAS and pneumococcus. Preferably, the GBS polynucleotide is selected from GBS Subset 1. In an alternative embodiment, the polynucleotide is a GBS polynucleotide which is homologous to at least one gene from both GAS and pneumococcus, but which is not homologous to a gene in any other published bacterial genome at the time of the invention. Preferably, the GBS polynucleotide is selected from one of the 12 GBS genes included in GBS Subset 1(a). (Table 3).

In another embodiment, the polynucleotide is a GAS polynucleotide which is homologous to at least one gene in both GBS and pneumococcus. Preferably, the GAS polynucleotide is selected from GAS Subset 1. In another embodiment, the polynucleotide is a GAS polynucleotide which is homologous to at least one gene in both GBS and pneumococcus but which is not homologous to any gene in any other published bacterial genome at the time of the invention. Preferably, the GAS polynucleotide is selected from GAS Subset 1(a).

Alternatively, the polynucleotide is a pneumoccus polynucleotide which is homologous to at least one gene in both GBS and GAS. Preferably, the pneumococcus polynucleotide is selected from Spn Subset 1(a). In another embodiment, the polynucleotide is a pneumoccus polynucleotide which is homologous to at least one gene in both GBS and GAS but which does not have a homologue in any other published bacterial genome at the time of the invention. Preferably, the pneumococcus polynucleotide is selected from Spn Subset 1(a).

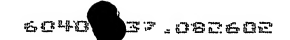
The invention further provides an immunogenic composition comprising a polypeptide, or a fragment or derivative thereof, which is encoded by a polynucleotide sequence which is specific to GBS. In one embodiment, the polynucleotide is a GBS polynucleotide which is not homologue to a gene in either GAS or pneumococcus. Preferably, the GBS polynucleotide is selected from one of the 683 GBS genes included in GBS Subset 4. In a further embodiment, the polynucleotide is a GBS polynucleotide which is not homologous to a gene in either GAS or pneumococcus or

any other published bacterial genome at the time of the invention. Preferably, the GBS polynucleotide is selected from one of the 315 GBS genes in GBS Subset 4(a).

The invention further provides an immunogenic composition comprising a polypeptide, or a fragment or derivative thereof, which is encoded by a polynucleotide sequence which is specific to GAS. In one embodiment, the polynucleotide is a GAS polynucleotide which is not homologous to a gene in either GBS or pneumococcus. Preferably, the GBS polynucleotide is selected from one of the 416 GAS genes included in GAS Subset 4. In a further embodiment, the polynucleotide is a GAS polynucleotide which does not have a homologue in either GBS or pneumococcus or in any other published bacterial genome at the time of the invention. Preferably, the GAS polynucleotide is selected from GAS Subset 4(a).

The invention further provides an immunogenic composition comprising a polypeptide, or a fragment or derivative thereof, which is encoded by a polynucleotide sequence which is specific to pneumococcus. In one embodiment, the polynucleotide is a pneumococcus polynucleotide which is not homologous to a gene in either GBS or GAS. Preferably, the pneumococcus polynucleotide is selected from one of the 836 Spn genes included in Spn Subset 4. In a further embodiment, the polynucleotide is a pneumococcus polynucleotide which does not have a homologue in either GBS or GAS or in any other published bacterial genome at the time of the invention. Preferably, the pneumococcus polynucleotide is selected from Spn Subset 4(a).

The invention further provides an immunogenic composition comprising a polypeptide, or a fragment or derivative thereof, which is encoded by a polynucleotide sequence which is specific to GBS and GAS. In one embodiment, the polynucleotide is a GBS polynucleotide which is homologous to at least one gene from GAS but is not homologous to a gene from pneumococcus. Preferably, the GBS polynucleotide is selected from one of the 225 GBS genes included in GBS Subset 2. In another embodiment, the GBS polynucleotide is homologous to at least one gene from GAS but is not homologous to any gene from pneumococcus and does not have a



homologue in any other published bacterial genome at the time of the invention. Preferably, the GBS polynucleotide is selected from GBS Subset 2(a).

In another embodiment, the polynucleotide is a GAS polynucleotide which is homologous to at least one gene from GBS but is not homologous to any gene from pneumococcus. Preferably, the GAS polynucleotide is selected from one of the 212 GAS genes included in GAS Subset 2. In another embodiment, the GAS polynucleotide is homologous to at least one gene from GBS but is not homologous to any gene from pneumococcus and does not have a homologous gene with any other published bacterial genome at the time of the invention. Preferably, the GAS polynucleotide is a selected from GAS Subset 2(a).

The invention further provides an immunogenic composition comprising a polypeptide, or a fragment or derivative thereof, which is encoded by a polynucleotide sequence which is specific to GBS and pneumococcus. In one embodiment, the polynucleotide is a GBS polynucleotide which is homologous to at least one gene from pneumococcus but is not homologous to any gene from GAS. Preferably, the GBS polynucleotide is selected from one of the 176 GBS genes included in GBS Subset 3. In another embodiment, the polynucleotide is a GBS polynucleotide which is homologous with at least one gene from pneumococcus but is not homologous with any GAS polynucleotide and does not have a homologous gene in any of the other published bacterial genomes at the time of the invention. Preferably, the GBS polynucleotide is selected from GBS Subset 3(a).

In another embodiment, the polynucleotide is a pneumococcus polynucleotide which is homologous with at least one gene from GBS, but is not homologous with any gene from GAS. Preferably, the pneumoccous polynucleotide is selected from one of the 195 Spn genes included in Spn Subset 2. In another embodiment, the polynucleotide is a pneumococcus polynucleotide which is homologous with at least one gene from GBS, but is not homologous with any gene from GAS and does not have a homologous gene in any other published bacterial genome at the time of the

invention. Preferably, the pneumococcus polynucleotide is selected from Spn Subset 3(a).

The invention further provides an immunogenic composition comprising a polypeptide, or a fragment or derivative thereof which is encoded by a polynucleotide sequence which is specific to GAS and pneumococcus. In one embodiment, the polynucleotide is a GAS polynucleotide which is homologous with at least one gene from pneumococcus but is not homologous with any gene from GBS. Preferably, the GAS polynucleotide is selected from one of the 62 GAS genes included in GAS Subset 3. In another embodiment, the polynucleotide is a GAS polynucleotide which is homologous with at least one gene from pneumococcus but is not homologous with any gene from GBS and is not homologous with any gene of any published bacterial genome at the time of the invention. Preferably, the GAS polynucleotide is selected from GAS Subset 3(a).

In another embodiment, the polynucleotide is a pneumococcus polynucleotide which is homologous with at least one GAS polynucleotide, but is not homologous with any GBS gene. Preferably, the pneumococus polynucleotide is selected from one of the 74 Spn genes included in Spn Subset 3. In another embodiment, the polynucleotide is a pneumococcus polynucleotide which is homologous with at least one gene from GAS, but is not homologous with any gene from GBS or with a gene from any other published bacterial genome at the time of the invention. Preferably, the pneumococcus polynucleotide is selected from Spn Subset 3(a).

The invention further provides an immunogenic composition comprising a polypeptide, or a fragment or derivative thereof, which is encoded by a polynucleotide sequence which is specific to one or more Streptococcal species serotypes.

Preferably, the polynucleotide is specific to a Streptococcal species serotype selected from the Streptococcal species GBS, GAS and pneumococcus. More preferably, the polynucleotide is specific to one or more GBS serotypes selected from the group consisting of GBS serotype Ia, Ib, II, III, IV, V, VI, VII and VIII.

The invention further provides an immunogenic composition comprising a polypeptide, or a fragment or derivative thereof, which is encoded by a polynucleotide sequence which is conserved across one or more Streptococcal species serotypes. Preferably, the polynucleotide is specific to a Streptococcal species serotype selected from the Streptococcal species GBS, GAS and pneumococcus. More preferable, the polynucleotide is conserved across one or more GBS serotypes selected from the group consisting of GBS serotype Ia, Ib, II, III, IV, V, VI, VII and VIII.

The invention further provides an immunogenic composition comprising a polypeptide, or a fragment or derivative thereof, which is encoded by a polynucleotide sequence which is specific to one or more clinical isolates of a Streptococcal species. Preferably, the polynucleotide is specific to a Streptococcal species clinical isolate selected from the Streptococcal species GBS, GAS and pneumococcus. More preferably, the polynucleotide is specific to one or more GBS clinical isolates selected from the clinical isolates identified in Table 5. Still more preferably, the polynucleotide is specific to one or more GBS clinical isolates having one or more genes selected from the genes listed in Table 7.

In another embodiment, the polynucleotide is a GBS polynucleotide which is homologous to at least one gene from both GAS and pneumococcus and which varies among clinical isolates. In another embodiment, the polynucleotide is a GBS polynucleotide which is homologous to at least one gene from both GAS and pneumococcus and which is homologous with at least one gene from at least one of the clinical isolates identified in Table 5. In another embodiment, the polynucleotide is a GBS polynucleotide which is homologous to at least one gene from both GAS and pneumococcus and which is homologous with at least one gene from each of the clinical isolates identified in Table 5. Preferably, the polynucleotide is selected from one of the genes listed in Table 7.

In one embodiment, the polynucleotide is a GBS polynucleotide which is homologous to at least one gene from GAS and is not homologous to any gene from pneumococcus and which varies among clinical isolates. In another embodiment, the

polynucleotide is a GBS polynucleotide which is homologous to at least one gene from GAS and is not homologous to any gene from pneumococcus and which is homologous to at least one gene from at least one of the clinical isolates identified in Table 5. In another embodiment, the polynucleotide is a GBS polynucleotide which is homologous to at least one gene from GAS and is not homologous to any gene from pneumococcus and which is homologous to at least one gene from each of the clinical isolates identified in Table 5. Preferably, the polynucleotide is selected from one of the genes listed in Table 7.

In one embodiment, the polynucleotide is a GBS polynucleotide which is homologous to at least one gene from pneumococcus and is not homologous to any gene from GAS and which varies among clinical isolates. In another embodiment, the polynucleotide is a GBS polynucleotide which is homologous to at least one gene from pneumococcus and is not homologous to any gene from GAS and which is homologous to at least one gene from at least one of the clinical isolates identified in Table 5. In another embodiment, the polynucleotide is a GBS polynucleotide which is homologous to at least one gene from pneumococcus and is not homologous to any gene from GAS and which is homologous to at least one gene from each of the clinical isolates identified in Table 5. Preferably, the polynucleotide is selected from one of the genes listed in Table 7.

In one embodiment, the polynucleotide is a GBS polynucleotide which is not homologous to any gene from GAS or pneumococcus and which varies among clinical isolates. In another embodiment, the polynucleotide is a GBS polynucleotide which is not homologous to any gene from GAS or pneumococcus and which is homologous to at least one gene from at least one of the clinical isolates identified in Table 5. In another embodiment, the polynucleotide is a GBS polynucleotide which is not homologous to any gene from GAS or pneumococcus and which is homologous to at least one gene from each of the clinical isolates identified in Table 5. Preferably, the polynucleotide is selected from one of the genes listed in Table 7.

The invention further provides an immunogenic composition comprising a polypeptide, or a fragment or derivative thereof, which is encoded by a polynucleotide sequence which is conserved across one or more clinical isolates of a Streptococcal species. Preferably, the polynucleotide is conserved across one or more Streptococcal clinical isolates selected from the Streptococcal species GBS, GAS and pneumococcus. More preferable, the polynucleotide is conserved across one or more GBS clinical isolates identified in Table 5. Still more preferably, the polynucleotide is conserved across one or more clinical isolates having one or more genes selected from the genes listed in Table 7.

The invention further provides for an immunogenic composition comprising a polypeptide encoded by a polynucleotide selected from one or more of the Subsets of the invention.

The invention provides a method for raising an immune response in a patient by administering any one of the immunogenic compositions set forth above. The choice of immunogenic composition means that the immune response may be reactive against all three of GAS, GBS and streptococcus, may be reactive against only two of the three, or may be reactive only against GBS.

The immune response is preferably an antibody response. It may be a protective immune response. The patient is preferably a human.

#### Essential genes and knockouts

The invention provides a *Streptococcus* bacterium wherein one or more genes within any of the Subsets of this invention have been knocked out. The choice of Subset means that the knocked out gene may be, for instance, a gene found in GBS but not in GAS or pneumococcus (e.g. which is involved in the pathogenesis of GBS, but not in the pathogenesis of GAS or pneumococcus, such as binding GBS cellular targets).

Techniques for producing knockout bacteria are well known, and knockout Streptococci of various species have been reported [e.g. Margolis et al. (2001)

Antimicrob. Agents Chemother. 45:2432-2435; Zhang et al. (2000) Cell 102:827-837; Nizet et al. (2000) Infect. Immun. 68:4245-4254; Nizet et al. (1997) Adv. Exp. Med. Biol. 418:627-630; etc.].

The knockout mutation may be situated in the coding region of the gene or may lie within its transcriptional control regions (e.g. within its promoter).

The knockout mutation will reduce the level of mRNA encoding the corresponding polypeptide to <1% of that produced by the wild-type bacterium, preferably <0.5%, more preferably <0.1%, and most preferably to 0%.

The knockout mutants of the invention may be used as immunogenic compositions (e.g. as vaccines) to prevent streptococcal infection. Such a vaccine may include the mutant as a live attenuated bacterium.

The knockout mutants of the invention may be used to determine whether genes are essential for bacterial survival, either under normal or stress conditions.

#### **Antisense**

The invention provides a single-stranded nucleic acid comprising a fragment of  $x_I$  or more nucleotides from a nucleotide sequence selected from one of the Subsets of the invention. The choice of group means that the nucleic acid may be complementary to a gene sequence found in GBS, GAS and pneumococcus, or a gene sequence specific to GBS.

The single-stranded nucleic acid is at least  $x_I$  nucleotides long. The value of  $x_I$  is at least 7 (e.g. 8, 9, 10, 11, 12, 13, 14, 15, 16, 17, 18, 19, 20, 21, 22, 23, 24, 25, 26, 27, 28, 29, 30, 31, 32, 33, 34, 35, 36, 37, 38, 39, 40, 41, 42, 43, 44, 45, 46, 47, 48, 49, 50 etc.). The single-stranded nucleic acid may be at most  $x_2$  nucleotides long, wherein  $x_2$  is 100 or less (e.g. 99, 98, 97, 96, 95, 94, 93, 92, 91, 90, 89, 88, 87, 86, 85, 84, 83, 82, 81, 80, 79, 78, 77, 76, 75, 74, 73, 72, 71, 70, 69, 68, 67, 66, 65, 64, 63, 62, 61, 60).

The nucleic acid is preferably of the formula 5'-(N)<sub>a</sub>-(X)-(N)<sub>b</sub>-3', wherein  $0\ge a\ge 15$ ,  $0\ge b\ge 15$ , N is any nucleotide, and X is the fragment as defined above. The

values of a and b may independently be 0, 1, 2, 3, 4, 5, 6, 7, 8, 9, 10, 11, 12, 13, 14 or 15. Each individual nucleotide N in the  $-(N)_{a}$  and  $-(N)_{b}$  portions of the nucleic acid may be the same or different. The length of the nucleic acid (i.e.  $a+b+x_{l}$ ) is preferably  $x_{2}$  or less.

Antisense inhibition of streptococcal gene expression is known e.g. Sato et al. (1998) FEMS Microbiol Lett 159:241-245. Antibacterial antisense techniques are also disclosed in international patent applications WO99/02673 and WO99/13893.

The single-stranded nucleic acid may reduce the level of polypeptide expression from the complementary gene to <1% of that produced by the wild-type bacterium, preferably <0.5%, more preferably <0.1%, and most preferably to 0%.

Antisense experiments may be used to determine whether genes are essential for bacterial survival, either under normal or stress conditions.

#### Screening methods

The invention provides a method for screening compounds, wherein the method involves contacting the compounds with a polypeptide expressed by one or more of the polynucleotides selected from one of the Subsets of the invention. The method may be for screening for agonists of the polypeptides, antagonists, antibiotics etc. The choice of group means, for instance, that the method may be used for identifying an antibiotic with broad anti-streptococcal activity could be identified, or for identifying an antibiotic specific to GBS.

Potential compounds for screening include small organic molecules, peptides, peptides, polypeptides, lipids, metals, nucleotides, nucleosides, aptamers, polyamines, antibodies, and derivatives thereof. Small organic molecules have a molecular weight between 50 and about 2,500 daltons, and most preferably in the range 200-800 daltons. Complex mixtures of substances, such as extracts containing natural products, compound libraries or the products of mixed combinatorial syntheses also contain potential antagonists.

Typically, a polypeptide is incubated with a test compound, and the mixture is then tested to see if the polypeptide and test compound interact, or to see if the polypeptide's activity is inhibited.

For preferred high-throughput screening methods, all the biochemical steps for this assay are performed in a single solution in, for instance, a test tube or microtitre plate, and the test compounds are analysed initially at a single compound concentration. For the purposes of high throughput screening, the experimental conditions are adjusted to achieve a proportion of test compounds identified as "positive" compounds from amongst the total compounds screened.

The invention also provides a compound identified using these methods. These can be used to treat or prevent streptococcal infection. The compound preferably has an affinity for the adhesion-specific protein of at least  $10^{-7}$  M e.g.  $10^{-8}$  M,  $10^{-9}$  M,  $10^{-10}$  M or tighter.

#### Distinguishing Streptococcal species

The invention provides a method for determining whether a Streptococcus bacterium of interest is or is not in the species agalactiae, pyogenes or pneumoiae, comprising the step(s) of: (a) contacting the bacterium with a nucleic acid probe comprising the sequence of a gene selected from one of the Subsets of the invention; and/or (b) contacting the bacterium with an antibody which binds to a polypeptide encoded by one or more of the polynucleotides of one or more of the Subsets of the invention. The choice of group means, for instance, that the method may be used for distinguishing GBS from GAS and from pneumococcus, or for confirming that a bacterium is not a GAS or pneumococcus.

The method will typically include the further step of detecting the presence or absence of an interaction between the bacterium of interest and the nucleic acid or protein.

The bacterium of interest may be in a cell culture, for example, or may be within a biological sample believed or known to contain a streptococcus. It may be intact or may be, for instance, lysed.

The term "biological sample" encompasses a variety of sample types obtained from an organism and can be used in a diagnostic or monitoring assay. The term encompasses blood and other liquid samples of biological origin, solid tissue samples, such as a biopsy specimen or tissue cultures or cells derived therefrom and the progeny thereof. The term encompasses samples that have been manipulated in any way after their procurement, such as by treatment with reagents, solubilization, or enrichment for certain components. The term encompasses a clinical sample, and also includes cells in cell culture, cell supernatants, cell lysates, serum, plasma, biological fluids, and tissue samples.

#### GBS 2603 Type V Genomic Sequence

Applicants have sequenced the complete genome sequence of GBS clinical type V isolate 2603 V/R and performed comparative analyses comparing this sequence with other GBS strains, with other species of pathogenic Streptococci and with other known bacterial species. The entire genomic sequence is available as of the filing date of this application at <a href="http://www.tigr.org">http://www.tigr.org</a>. This genomic sequence is incorporated herein by reference in its entirety. The genomic sequence of GBS type V isolate 2603 V/R is also set forth in International Patent Application WO 02/34771.

In one embodiment, the invention relates to the polynucleotides, and fragments and derivatives thereof, set forth in the GBS clinical type V isolate 2603 which are not disclosed within WO 02/34771. The invention further relates to polypeptides expressed by the polynucleotides of the invention.

Applicants have predicted that the GBS 2603 isolate contains approximately 2,176 predicted genes. Each predicted gene is set forth in Table 1, listed by a SAGxxxx ORF number. Table 1 also includes the predicted amino acid size of the

predicted expressed protein and the predicted function, if known. The sequence of each SAG reference can be obtained at the TIGR website.

Figure 1 is a circular representation of the GBS genome and comparative hybridisations using microarrays. The outer circle represents predicted coding regions on the plus strand color coded by role categories: violet indicating amino acid biosynthesis; light blue indicating biosynthesis of cofactors, prosthetic groups, and carriers; light green indicating cell envelope; red indicating cellular processes; brown indicating central intermediary metabolism; yellow indicating DNA metabolism; light gray indicating energy metabolism; magenta indicating fatty acid and phospholipid metabolism; pink indicating protein synthesis and fate; orange indicating purines, pyrimidines, nucleosides, and nucleotides; olive indicating regulatory functions and signal transduction; dark green indicating transcription; teal indicating transport and binding proteins; gray indicating unknown function; salmon indicating other categories; blue indicating hypothetical proteins.

The second circle represents predicted coding regions on the minus strand. In the third circle, black represents atypical nucleotide composition curve; green represents most atypical regions; magenta represents insertion elements; red diamonds indicate rRNAs.

Circles 4-22 represent comparative hybridisations of strain 2603 V/R with 19 GBS strains. Cy3/Cy5 (2603 V/R signal/test strain) ratio cutoffs were defined arbitrarily as Cy3/Cy5 -1.0-3.0, the gene was present in the test strain, no color was added; Cy3/Cy5 = 3.0-10.0, ambiguous result (blue); Cy3/Cy5 > 10, gene absent in test strain (red).

Circles 4 – 9 represent type 1a strains 090, 515, A909, Davis, and DK8.

Circles 10 – 11 represent type 1b strains S7 7357b and H36B. Circles 12 – 13 represent type II strains 18RS21 and DK21. Circles 14 – 18 represent type III COH1, COH31, D136C, M732 and M781. Circle 19 represents type V strain CJB111.

Circles 20 – 21 represent type VIII strains SMU014 and JM9130013. Circle 22

represents nontypable (NT) strain CJB110. Throughout Figure 1, varying regions of five or more consecutive genes are indicated by yellow bullets.

Figure 4 depicts a linear representation of the GBS genome. The location of predicted coding regions color-coded by biological role (see Figure 1) is displayed. Arrowed boxes represent the direction of transcription for each ORF. The number of membrane-spanning domains predicted by TopPred is displayed as lipid bi-layers on top of ORFs, only for those whose products have five or more predicted membrane spanning regions. Genes coding for rRNAs (16S, 23S, 5S) and tRNAs (clover leaf structure with number of genes) are indicated. Predicted Rho-independent transcriptional terminators are represented by hairpins.

ORF's were predicted by GLIMMER (See, Delcher, et al., (1999) Nucleic Acids Res. 27, 4636 - 4641 and Salzberg, et al., (1998) Nucleic Acids Res. 26, 544-548) trained with ORFs larger than 600 base pairs from the genomic sequence and GBS genes available in GenBank. All predicted proteins larger than 30 amino acids were searched against a nonredundant protein database. (See Fleischmann, et al., (1995) Science 269, 496 - 512). Frame-shifts and point mutations were detected and corrected where appropriate; those remaining were annotated as "authentic frameshift" or "authentic point mutation". Protein membrane-spanning domains were identified by TOPPRED (See Claros, et al., (1994) Comput. Appl. Biosci. 10, 685 -686). Candidate lipoprotein signal peptides (See Hayashi et al., (1990) J. Bioenerg. Biomembr. 22, 451 - 471) were flagged by N-terminal exact matches to the pattern {DERK} (6)-[LIVMFWSTAG] (2)-[LIVMFYSTAGCQ] - [AGS] - C. Putative signal peptides were identified by using SIGNALP (Nielsen, et al., (1997) Protein Eng. 10, 1 - 6). Two sets of hidden Markov models were used to determine ORF membership in families and superfamilies: PFAM Ver. 5.5 (Bateman, et al., (2000) Nucleic Acids Res. 28, 263 - 266) and TIGRFAMS 1.0 (Haft et al., (2001) Nucleic Acids Res. 29, 41 - 43). Domain-based paralogous families were built by performing all-versus-all searches on the protein sequences by using a modified version of a previously described method. (Niermann, et al., (2001) Proc. Natl. Acad. Sci. USA

98, 4136 - 4141) Potential lineage-specific gene duplications were estimated by identification of OFRs more similar to ORFs within the GBS genome than to ORFs from other complete genomes. All ORFs were searched with FASTA3 (Pearson (2000) *Methods Mol. Biol.* 132, 185 - 219) against all ORF's from the complete genomes and matches with a FASTA P value of 10<sup>-15</sup> were considered significant.

The genome consists of a circular chromosome of 2,160,266 base pairs with a G+C content of 35.7%. Base pair one of the chromosome was assigned within the putative origin of replication. The genome contains 80 tRNAs, 7rRNAs, and 3 sRNAs. Approximately 78% of the 2,176 predicted genes are transcribed in the same direction as that of DNA replication, a feature also observed in S. pn. and other low-GC Gram positive organisms.

Biological roles were assigned to 1,409 (65%) of the genome according to a classification scheme adapted from Riley (1993) *Microbiol. Rev.* 57, 862 - 952. Another 527 predicted proteins (24%) matched proteins of unknown function, and the remaining 240 (11%) had no database match. The expression of 50 of these hypothetical proteins was confirmed by Western Blot analysis, and the proteins were annotated as "proteins of unknown function." A total of 339 paralogous protein families were identified in strain 2603, containing 941 predicted proteins (43% of the total).

The Western Blot analysis was conducted as follows. GBS strain 2603 V/R cells were grown in Todd-Hewitt broth (Difco) to OD600nm = 0.5. The culture was centrifuged for 20 minutes at 5,000 rpm. The supernatant was discarded, and bacteria were washed once with PBS, resuspended in 2 ml of 50 mM Tris-HCl pH 6.8, containing 400 units of Mutanolysin (Sigma), and incubated 2 hours at 37°C. After three cycles of freeze and thaw, cellular debris was removed by centrifugation at 14,000 rpm for 10 minutes, and the protein concentration of the supernatant was measured by the Bio-Rad Protein assay, with BSA as a standard. Purified recombinant proteins (50 ng) and total cell extracts (25 µg) derived from GBS serotype V 2603 V/R strain were separated by SDS/PADE and electroblotted onto

nitrocellulose membranes for 1 hour at 100 V. The membranes were saturated by overnight incubation at 4° C in 5% skimmed milk and 0.1% Tween 20 in PBS and incubated for 1 hour at room temperature with sera from immunized mice diluted 1:500 - 1:1,000 in saturation buffer. To reduce background due to antibodies raised against contaminating *E. coli* proteins, sera were preincubated with *E. coli* protein extracts absorbed on nitrocellulose strips. The membranes were washed twice in 3% skimmed milk and 0.1% Tween 20 in PBS and incubated for 1 hour with a 1:1,000 dilution of horseradish peroxidase-conjugated antimouse Ig (DAKO). After washing with 0.1% Tween 20 in PBS, the membranes were developed with the Opti-4CN Substrate Kit (Bio-Rad).

Table 2 comprises a list of predicted and experimentally characterized surface and secreted proteins from GBS. Candidate signal peptides and lipoprotein motifs were predicted with PSORT [Nakai, K. & Horton, P. (1999) Trends Biochem Sci 24, 34-6] and other methods (see methods), sortase motifs (LPxTG) were detected using the FINDPATTERNS program of the GCG Package [Devereux, J., Haeberli, P. & Smithies, O. (1984) Nucleic Acids Res 12, 387-95] and hidden Markov models. Column "Other" indicates proteins carrying other motifs (e.g. integrin-binding motif RGD) or are similar to characterized surface-exposed proteins. Western blot results were considered positive when the antibodies revealed a predominant band of the expected molecular weight on the total protein extracts of S. agalactiae strain 2603 V/R, ORFs without + or - in this column were not tested in western blot. FACS analyses were performed for western blot positive proteins only. Western blot and FACS data are displayed only for proteins carrying at least one of the other motifs shown in the table. Column "GBS specific" indicates genes unique to S. agalactiae (when compared to other completely sequenced genomes) that are present in all the S. agalactiae strains tested in comparative genome hybridization analyses. Finally, only proteins carrying less than 3 predicted transmembrane domains are shown in the table, other proteins are likely to be embedded in the cytoplasmic membrane and are probably not exposed on the organism's surface.

FACS data was collected as follows: GBS 2603 V/R strain cells were grown in Todd-Hewitt broth (Difco) to OD600nm = 0.5. The culture was centrifuged for 20 minutes at 5,000 rpm, and bacteria were washed once with PBS, resuspended in PBS containing 0.05% paraformaldehyde, and incubated for 1 hour at 37°C and then overnight at 4°C. Fifty microliters of fixed bacteria (OD600nm 0.1) was washed once with PBS, resuspended in 20  $\mu$ I of newborn calf serum (Sigma), and incubated for 1 hour at 4°C in 100µI of preimmune or immune sera and diluted 1:200 in dilution buffer (PBS, 20% newborn calf serum, 0.1% BSA). After centrifugation and washing with 200µl of washing buffer (0.1% BSA in PBS), samples were incubated for 1 hour at 4°C with 50 µl of R-phycoerythrin-conjugated F(ab)2 goat anti-mouse IgG (Jackson ImmunoResearch) diluted 1:100 in dilution buffer. Cells were washed with 200  $\mu l$  of washing buffer and resuspended in 200  $\mu l$  of PBS. Samples were analysed by using a FACS calibur apparatus (Becton Dickinson), and data were analyzed by using CELL QUEST (Becton Dickinson). A shift in mean fluorescence intensity of >75 channels compared with preimmune sera from the same mice was considered positive. This cutoff was determined from the mean plus two standard deviations of shifts obtained with control sera raised against mock purified recombinant proteins from cultures of E. coli carrying the empty expression vector and included in every experiment. Artifacts due to bacterial lysis were excluded by using antisera raised against six different known cytoplasmic proteins, all of which gave negative results.

# Regions of Atypical Nucleotide Composition.

These regions were identified by the  $x^2$  analysis: the distribution of all 64 trinucleotides (3 mers) was computed for the complete genome in all six reading frames, followed by the 3-mer distribution in 2,000-bp windows. Windows overlapped by 1,000 bp. For each window, the  $x^2$  statistic on the difference between its 3-mer content, and that of the whole genome was computed.

## In Silico Genome Comparisons

The protein sets of S. agalactiae, Streptococcus pneumoniae and S. pyogenes were compared by using FASTA3. A general description of the FASTA3 sequence comparison program is discussed in Pearson, W.R., "Flexible Sequence Similarity Searching with the FASTA3 Program Package", (2000) Methods Mol. Biol., 132: 185-219. Shared genes were defined using a FASTA3 P value cutoff of 10<sup>-15</sup>. These shared genes and genes that S. agalactiae did not share with the other streptococci using this cutoff were subsequently searched against all completely sequenced genomes, and genes were defined as unique to streptococci or S. agalactiae when they did not share similarity with any other gene sets with a FASTA3 P value of 10<sup>-5</sup> or lower. The use of two cutoffs provides for a more stringent analysis of shared or unique genes.

Figure 2 is a schematic representation of in silico comparisons between streptococci. The protein sets of GBS, S. pn., and GAS were compared by using FASTA3. Numbers under the species name indicate genes that are not shared with the other species; values in parenthesis are the number of proteins in each species (excluding frame-shifted and degenerated genes). Numbers in the intersections indicate genes shared by two or three species. These are displayed in the color corresponding to the species used as the query. (GBS: green; S.pn.: blue; GAS: red). Numbers in any given intersection are slightly different due to gene duplications in some species.

Table 3 lists genes which were shared among GBS, GAS and pneumococcus, but which were not found in any of the other completely sequenced genomes. The protein sets of S. agalactiae, S. pneumoniae, and S. pyogenes were compared using FASTA3 [Pearson, W. R. (2000) Methods Mol Biol 132, 185-219]. Shared genes were defined using a FASTA3 p value cutoff of 10<sup>-15</sup>. These shared genes and genes that S. agalactiae did not share with the other streptococci using this cutoff were subsequently searched against all completely sequenced genomes and genes were

defined as unique to streptococci or *S. agalactiae* when they did not share similarity with any other gene sets with a FASTA3 p value of 10<sup>-5</sup> or lower.

### **Synteny**

Regions of conservation of gene synteny were computed as windows of 10 kb spanning at least three genes whose order was conserved in the other species.

Regions were merged if they were less than 20 kb apart. The number of genes within each broad region was then calculated.

#### Comparative Genome Hybridizations

Comparative genome hybridizations (See Figure 1) using DNA microarrays were performed between the sequenced type V strain 2603 V/R and 19 other GBS strains of multiple serotypes (See Table %). Predicted genes from strain 2603 V/R were amplified by PCR and arrayed on glass microscope slides. See Peterson, et al., (2000) J. Bacteriol. 182, 6192-6202. Genomic DNA was labelled according to protocols provided by J. DeRisi (www.microarrays.org/Pdfs/Genomic-<u>DNALabel B.pdf</u>), except that the DNA was not digested or sheared before labelling. Arrays were scanned with a GENEPIX 4000B scanner (Axon Instruments, Foster City, CA), and individual hybridisation signals were quantitated with TIGR SPOTFINDER. See Hedge, et al., (2000), Biotechniques 29, 548-550, 552-554, 556. Cy3/Cy5 (2603 V/R signal/test strain) ratio cutoffs were defined arbitrarily as Cy3/Cy5 = 1.0 - 3.0, gene present in test strain; 3.0 - 10.0, ambiguous result; >10.0. gene absent. For ambiguous results, the gene may be divergent in the test strain relative to 2603 V/R, or the gene may be absent in the test strain but still produces paralogous gene family or a repetitive elemtn. Although cutoffs are arbitrary, they fit nicely the results for the variation of the capsule locus in the strains tested (see region 9 on Figure 1) where most genes are slightly divergent and only a few are completely different.



The CGH detected 1,698 genes in all of the strains, whereas 401 genes from strain 2603 V/R (18% of the gene complement) were not detected in at least one other strain, suggesting that they are absent or significantly divergent in those strains. Two hundred sixty (38%) of the 683 genes specific to S. agalactiae when compared with the other two streptococci (Fig. 2), including virulence determinants and surface proteins, vary among S. agalactiae strains, whereas only 47 (4%) of the genes common to all three streptococcal species, including 5 of the 6 sortases identified in the genome, vary among strains. Thus, the in silico analysis of genes shared by the streptococci that are not expected to vary among this genus is consistent with the CGH analysis. Forty-four (25%) of the genes shared by S. agalactiae and S. pneumoniae and 44 (20%) of those shared by S. agalactiae and S. pyogenes vary in the CGH analysis. The first set contains many glycosyl transferases and proteins carrying a cell-wall anchor, whereas the second set displays many phage-related genes. One hundred thirty-six of the 315 genes unique to S. agalactiae when compared with all sequenced genomes vary among strains. These include R5, three capsular genes, two cell wall-anchored proteins, and three transcriptional regulators. Three hundred sixty-four (91%) of the 401 varying genes correspond to 15 regions containing more than 5 contiguous genes. Ten of these regions display an atypical nucleotide composition in strain 2603 V/R (Fig. 1), consistent with the possibility that they were horizontally transferred into this strain. Two of the largest regions (region 4, a prophage and region 7, similar to Tn916 from Enterococcus faecalis) are flanked by insertion sequence elements. The 15 regions contain many proteins predicted to be anchored on the cell wall or surface exposed, including Rib (region 3), sortases, glycosyl transferases, the capsule locus (region 9, divergent in all strains but the other type V strain CJB111), and phage-related genes. Region 14 is unique to S. agalactiae and spans 33 genes (SAG1989- SAG2021), including 25 proteins of unknown function, some of which carry a cell-wall anchor. It is flanked by an ISL3 transposase and displays an atypical nucleotide composition. Region 1, unique to S. agalactiae, is a possible plasmid or remnant of a phage (SAG0218-SAG0238), contains mostly

hypothetical proteins, and is flanked by a site-specific recombinase. Region 8 is specific to *S. agalactiae*, comprises 20 proteins of unknown function (SAG1018-SAG1037), most of which are predicted to be membrane associated or secreted, and displays an atypical nucleotide composition.

The CGHresults were analyzed by profile clustering where genes are grouped based on their distribution patterns (Fig. 5). Sixteen clusters of five or more contiguous and noncontiguous genes comprising a total of 300 genes were identified (Table 6). Several clusters correspond to regions of contiguous genes described above. Some clusters of genes that do not share sequence similarity and are located at different loci in the genome display an identical profile. For instance, a cluster of genes containing a surface antigen (SAG0674-SAG0681) follows the same distribution as another cluster containing only hypothetical proteins (SAG0247-SAG0249). A putative pathogenicity protein (SAG2063) also clusters with a region containing several glycosyl transferases and Sec proteins (SAG1447-SAG1462).

Profile clustering was also used to group strains based on similarity of gene content (Fig. 5). In addition, the sequences of 19 genes from each of 11 *S. agalactiae* strains were determined after PCR amplification and used for phylogenetic analyses. The strains were the following: type Ia, 090 and A909; type Ib, H36B; type II, 18RS21; type III, COH1, M732 and M781; type V, 2603 V/R and 1169NT1; type VIII, JM9130013; and nontypeable strain CJB110. The set comprised 8 housekeeping genes and 11 genes coding for proteins predicted to be surface-exposed (Table 7).

The profile clustering was conducted as follows. The information and absence of genes based on the comparative genome hybridisation results was used to group genes based on their distribution patterns. The analysis used was essentially identical to that used for phylogenetic profile analysis. See Pellegrinie, et al., (1999) *Proc.*Natl. Acad. Sci. USA 96, 4285 – 4288. Each gene was assigned a binary profile based on its presence or absence across the different strains, with presence determined by a

Cy3/Cy5 ratio < 3.0 and absence ≥ 3.0. The gene profiles were then clustered by using the single-linkage clustering algorithm with column weighting (all with default settings) of CLUSTER (<a href="http://rana.lbl.gov">http://rana.lbl.gov</a>). The CLUSTER program also groups the strains (columns) based on similarity of gene profiles. Clusters of genes and strains were viewed by using TREEVIEW (<a href="http://rana.lbl.gov">http://rana.lbl.gov</a>).

Phylogenetic trees were inferred for the complete set of 19 genes and for the subsets of housekeeping and surface-exposed genes. Because the branching patterns in all three trees were identical, only the tree of the 19 genes is shown in Fig. 3. The degree of polymorphism of the housekeeping and the surface-exposed genes is similar (~1 variable site among all of the strains per 100 bp).

The sequences of genes from the different strains were aligned by using CLUSTALW (See Thompson (1994), *Nucleic Acids Res.* 22, 4673 – 4680.) and trimmed to remove ambiguously aligned regions. Phylognetic trees of individual genes and of concatenated alignments of multiple genes were inferred by using maximum likelihood methods of PAUP\* 4.0 b10 (Sinauer, Sunderland, MA). Bootstrap analysis was carried out using PAUP\* as well. The possibility of recombination among strains was examined by using analysis of sequence variation using SIMPLOT (S.C. Ray) and analysis of phylogenetic heterogeneity by using MACCLADE (Sinauer).

Analysis of this variation showed no evidence for major recombination events between the strains. There were no long stretches of polymorphic sites that strongly supported other trees (analysis with MACCLADE), and there were no significant crossover events in plots of sequence similarity between strains (analysis with SIMPLOT). Some strain groupings (clades) generated by phylogenetic analysis were similar to clusters from the profile analysis (type III strains M781, M732 and COH1; type Ia strain 090 and nontypable strain CJB110), whereas others were different, possibly because of the aforementioned problems with the profile clustering. In both the phylogenetic analysis and the profile clustering, there is serotypedependent and -

independent clustering (Figs. 3 and 5). The presence of strains of the same serotype in different clades or clusters could be due to lateral gene transfer.

Figure 5 demonstrates phylogenetic profiling of GBS strains based on comparative genome hybridisations. The information on presence and absence of genes based on the microarray comparative genome hybridization results was used for phylogenetic profile analysis. The presence of a particular gene or gene cluster is indicated in the figure by a red square and the absence of a gene or cluster by a black square. The relationship between strains based on this analysis is depicted by the tree at the top of the figure. The strains and their serotypes are indicated (NT: nontypeable). Clusters with identical profiles are reduced to a single horizontal line and the number of genes in each cluster is indicated on the right. The clusters of 5 or more genes, labeled in red text and numbered, are listed in Table 6. The 1698 genes shared by all 19 strains are labeled in green text.

Figure 3 depicts a phylogenetic tree of GBS strains based on PCR sequences. The sequences of 19 genes (Table 7) from each of 11 GBS strains were aligned and trimmed to remove ambiguously aligned regions, and phylogenetic trees were inferred. Strain names are indicated in bold, and serotypes are indicated under the strain names. Bootstrap values are indicated on the branches.

#### **Techniques**

A summary of standard techniques and procedures which may be employed in order to perform the invention (e.g. to utilise the disclosed sequences for vaccination or diagnostic purposes) follows. This summary is not a limitation on the invention, but gives examples that may be used, but are not required.

#### **General**

The practice of the present invention will employ, unless otherwise indicated, conventional techniques of molecular biology, microbiology, recombinant DNA, and immunology, which are within the skill of the art. Such techniques are explained fully in the

literature eg. Sambrook Molecular Cloning; A Laboratory Manual, Second Edition (1989) or Third Edition (2000); DNA Cloning, Volumes I and II (D.N Glover ed. 1985); Oligonucleotide Synthesis (M.J. Gait ed, 1984); Nucleic Acid Hybridization (B.D. Hames & S.J. Higgins eds. 1984); Transcription and Translation (B.D. Hames & S.J. Higgins eds. 1984); Animal Cell Culture (R.I. Freshney ed. 1986); Immobilized Cells and Enzymes (IRL Press, 1986); B. Perbal, A Practical Guide to Molecular Cloning (1984); the Methods in Enzymology series (Academic Press, Inc.), especially volumes 154 & 155; Gene Transfer Vectors for Mammalian Cells (J.H. Miller and M.P. Calos eds. 1987, Cold Spring Harbor Laboratory); Mayer and Walker, eds. (1987), Immunochemical Methods in Cell and Molecular Biology (Academic Press, London); Scopes, (1987) Protein Purification: Principles and Practice, Second Edition (Springer-Verlag, N.Y.), and Handbook of Experimental Immunology, Volumes I-IV (D.M. Weir and C. C. Blackwell eds 1986).

Standard abbreviations for nucleotides and amino acids are used in this specification.

#### Further Definitions

A composition containing X is "substantially free of" Y when at least 85% by weight of the total X+Y in the composition is X. Preferably, X comprises at least about 90% by weight of the total of X+Y in the composition, more preferably at least about 95% or even 99% by weight. The term "comprising" means "including" as well as "consisting" e.g. a composition "comprising" X may consist exclusively of X or may include something additional e.g. X + Y. The singular forms "a", "and", and "the" include plural referents unless the context clearly dictates otherwise. Thus, for example, reference to "a polynucleotide" includes a plurality of such polynucleotides and reference to "an epithelial cell" includes reference to one or more cells and equivalents thereof known to those skilled in the art, etc.

The term "heterologous" refers to two biological components that are not found together in nature. The components may be host cells, genes, or regulatory regions, such as promoters. Although the heterologous components are not found together in nature, they can function together, as when a promoter heterologous to a gene is operably linked to the gene. Another example is where a Streptococcal sequence is heterologous to a mouse host cell. A further

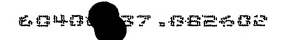
examples would be two epitopes from the same or different proteins which have been assembled in a single protein in an arrangement not found in nature.

An "origin of replication" is a polynucleotide sequence that initiates and regulates replication of polynucleotides, such as an expression vector. The origin of replication behaves as an autonomous unit of polynucleotide replication within a cell, capable of replication under its own control. An origin of replication may be needed for a vector to replicate in a particular host cell. With certain origins of replication, an expression vector can be reproduced at a high copy number in the presence of the appropriate proteins within the cell. Examples of origins are the autonomously replicating sequences, which are effective in yeast; and the viral T-antigen, effective in COS-7 cells.

A "mutant" sequence is defined as DNA, RNA or amino acid sequence differing from but having sequence identity with the native or disclosed sequence. Depending on the particular sequence, the degree of sequence identity between the native or disclosed sequence and the mutant sequence is preferably greater than 50% (eg. 60%, 70%, 80%, 90%, 95%, 99% or more, calculated using the Smith-Waterman algorithm as described above). As used herein, an "allelic variant" of a nucleic acid molecule, or region, for which nucleic acid sequence is provided herein is a nucleic acid molecule, or region, that occurs essentially at the same locus in the genome of another or second isolate, and that, due to natural variation caused by, for example, mutation or recombination, has a similar but not identical nucleic acid sequence. A coding region allelic variant typically encodes a protein having similar activity to that of the protein encoded by the gene to which it is being compared. An allelic variant can also comprise an alteration in the 5' or 3' untranslated regions of the gene, such as in regulatory control regions (eg. see US patent 5,753,235).

#### Expression systems

The Streptococcal nucleotide sequences can be expressed in a variety of different expression systems; for example those used with mammalian cells, baculoviruses, plants, bacteria, and yeast.



## i. Mammalian Systems

Mammalian expression systems are known in the art. A mammalian promoter is any DNA sequence capable of binding mammalian RNA polymerase and initiating the downstream (3') transcription of a coding sequence (eg. structural gene) into mRNA. A promoter will have a transcription initiating region, which is usually placed proximal to the 5' end of the coding sequence, and a TATA box, usually located 25-30 base pairs (bp) upstream of the transcription initiation site. The TATA box is thought to direct RNA polymerase II to begin RNA synthesis at the correct site. A mammalian promoter will also contain an upstream promoter element, usually located within 100 to 200 bp upstream of the TATA box. An upstream promoter element determines the rate at which transcription is initiated and can act in either orientation [Sambrook et al. (1989) "Expression of Cloned Genes in Mammalian Cells." In Molecular Cloning: A Laboratory Manual, 2nd ed.].

Mammalian viral genes are often highly expressed and have a broad host range; therefore sequences encoding mammalian viral genes provide particularly useful promoter sequences. Examples include the SV40 early promoter, mouse mammary tumor virus LTR promoter. adenovirus major late promoter (Ad MLP), and herpes simplex virus promoter. In addition, sequences derived from non-viral genes, such as the murine metallotheionein gene, also provide useful promoter sequences. Expression may be either constitutive or regulated (inducible), depending on the promoter can be induced with glucocorticoid in hormone-responsive cells. The presence of an enhancer element (enhancer), combined with the promoter elements described above, will usually increase expression levels. An enhancer is a regulatory DNA sequence that can stimulate transcription up to 1000-fold when linked to homologous or heterologous promoters, with synthesis beginning at the normal RNA start site. Enhancers are also active when they are placed upstream or downstream from the transcription initiation site, in either normal or flipped orientation, or at a distance of more than 1000 nucleotides from the promoter [Maniatis et al. (1987) Science 236:1237; Alberts et al. (1989) Molecular Biology of the Cell, 2nd ed.]. Enhancer elements derived from viruses may be particularly useful, because they usually have a broader host range. Examples include the SV40 early gene enhancer [Dijkema et al (1985) EMBO J. 4:761] and the enhancer/promoters derived from the long

terminal repeat (LTR) of the Rous Sarcoma Virus [Gorman et al. (1982b) *Proc. Natl. Acad. Sci.* 79:6777] and from human cytomegalovirus [Boshart et al. (1985) *Cell* 41:521]. Additionally, some enhancers are regulatable and become active only in the presence of an inducer, such as a hormone or metal ion [Sassone-Corsi and Borelli (1986) *Trends Genet.* 2:215; Maniatis et al. (1987) *Science* 236:1237].

A DNA molecule may be expressed intracellularly in mammalian cells. A promoter sequence may be directly linked with the DNA molecule, in which case the first amino acid at the N-terminus of the recombinant protein will always be a methionine, which is encoded by the ATG start codon. If desired, the N-terminus may be cleaved from the protein by *in vitro* incubation with cyanogen bromide.

Alternatively, foreign proteins can also be secreted from the cell into the growth media by creating chimeric DNA molecules that encode a fusion protein comprised of a leader sequence fragment that provides for secretion of the foreign protein in mammalian cells. Preferably, there are processing sites encoded between the leader fragment and the foreign gene that can be cleaved either *in vivo* or *in vitro*. The leader sequence fragment usually encodes a signal peptide comprised of hydrophobic amino acids which direct the secretion of the protein from the cell. The adenovirus triparite leader is an example of a leader sequence that provides for secretion of a foreign protein in mammalian cells.

Usually, transcription termination and polyadenylation sequences recognized by mammalian cells are regulatory regions located 3' to the translation stop codon and thus, together with the promoter elements, flank the coding sequence. The 3' terminus of the mature mRNA is formed by site-specific post-transcriptional cleavage and polyadenylation [Birnstiel et al. (1985) Cell 41:349; Proudfoot and Whitelaw (1988) "Termination and 3' end processing of eukaryotic RNA. In Transcription and splicing (ed. B.D. Hames and D.M. Glover); Proudfoot (1989) Trends Biochem. Sci. 14:105]. These sequences direct the transcription of an mRNA which can be translated into the polypeptide encoded by the DNA. Examples of transcription terminater/polyadenylation signals include those derived from SV40 [Sambrook et al (1989) "Expression of cloned genes in cultured mammalian cells." In Molecular Cloning: A Laboratory Manual].

Usually, the above described components, comprising a promoter, polyadenylation signal, and transcription termination sequence are put together into expression constructs. Enhancers, introns with functional splice donor and acceptor sites, and leader sequences may also be included in an expression construct, if desired. Expression constructs are often maintained in a replicon, such as an extrachromosomal element (eg. plasmids) capable of stable maintenance in a host, such as mammalian cells or bacteria. Mammalian replication systems include those derived from animal viruses, which require trans-acting factors to replicate. For example, plasmids containing the replication systems of papovaviruses, such as SV40 [Gluzman (1981) Cell 23:175] or polyomavirus, replicate to extremely high copy number in the presence of the appropriate viral T antigen. Additional examples of mammalian replicons include those derived from bovine papillomavirus and Epstein-Barr virus. Additionally, the replicon may have two replicaton systems, thus allowing it to be maintained, for example, in mammalian cells for expression and in a prokaryotic host for cloning and amplification. Examples of such mammalian-bacteria shuttle vectors include pMT2 [Kaufman et al. (1989) Mol. Cell. Biol. 9:946] and pHEBO [Shimizu et al. (1986) Mol. Cell. Biol. 6:1074].

The transformation procedure used depends upon the host to be transformed. Methods for introduction of heterologous polynucleotides into mammalian cells are known in the art and include dextran-mediated transfection, calcium phosphate precipitation, polybrene mediated transfection, protoplast fusion, electroporation, encapsulation of the polynucleotide(s) in liposomes, and direct microinjection of the DNA into nuclei.

Mammalian cell lines available as hosts for expression are known in the art and include many immortalized cell lines available from the American Type Culture Collection (ATCC), including but not limited to, Chinese hamster ovary (CHO) cells, HeLa cells, baby hamster kidney (BHK) cells, monkey kidney cells (COS), human hepatocellular carcinoma cells (eg. Hep G2), and a number of other cell lines.

### ii. Baculovirus Systems

The polynucleotide encoding the protein can also be inserted into a suitable insect expression vector, and is operably linked to the control elements within that vector. Vector construction

employs techniques which are known in the art. Generally, the components of the expression system include a transfer vector, usually a bacterial plasmid, which contains both a fragment of the baculovirus genome, and a convenient restriction site for insertion of the heterologous gene or genes to be expressed; a wild type baculovirus with a sequence homologous to the baculovirus-specific fragment in the transfer vector (this allows for the homologous recombination of the heterologous gene in to the baculovirus genome); and appropriate insect host cells and growth media.

After inserting the DNA sequence encoding the protein into the transfer vector, the vector and the wild type viral genome are transfected into an insect host cell where the vector and viral genome are allowed to recombine. The packaged recombinant virus is expressed and recombinant plaques are identified and purified. Materials and methods for baculovirus/insect cell expression systems are commercially available in kit form from, inter alia, Invitrogen, San Diego CA ("MaxBac" kit). These techniques are generally known to those skilled in the art and fully described in Summers & Smith, Texas Agricultural Experiment Station Bulletin No. 1555 (1987) ("Summers & Smith").

Prior to inserting the DNA sequence encoding the protein into the baculovirus genome, the above described components, comprising a promoter, leader (if desired), coding sequence, and transcription termination sequence, are usually assembled into an intermediate transplacement construct (transfer vector). This may contain a single gene and operably linked regulatory elements; multiple genes, each with its owned set of operably linked regulatory elements; or multiple genes, regulated by the same set of regulatory elements. Intermediate transplacement constructs are often maintained in a replicon, such as an extra-chromosomal element (e.g. plasmids) capable of stable maintenance in a host, such as a bacterium. The replicon will have a replication system, thus allowing it to be maintained in a suitable host for cloning and amplification.

Currently, the most commonly used transfer vector for introducing foreign genes into AcNPV is pAc373. Many other vectors, known to those of skill in the art, have also been designed. These include, for example, pVL985 (which alters the polyhedrin start codon from ATG to ATT, and

which introduces a BamHI cloning site 32 basepairs downstream from the ATT; see Luckow and Summers, Virology (1989) 17:31.

The plasmid usually also contains the polyhedrin polyadenylation signal (Miller et al. (1988) Ann. Rev. Microbiol., 42:177) and a prokaryotic ampicillin-resistance (amp) gene and origin of replication for selection and propagation in E. coli.

Baculovirus transfer vectors usually contain a baculovirus promoter. A baculovirus promoter is any DNA sequence capable of binding a baculovirus RNA polymerase and initiating the downstream (5' to 3') transcription of a coding sequence (eg. structural gene) into mRNA. A promoter will have a transcription initiation region which is usually placed proximal to the 5' end of the coding sequence. This transcription initiation region usually includes an RNA polymerase binding site and a transcription initiation site. A baculovirus transfer vector may also have a second domain called an enhancer, which, if present, is usually distal to the structural gene. Expression may be either regulated or constitutive.

Structural genes, abundantly transcribed at late times in a viral infection cycle, provide particularly useful promoter sequences. Examples include sequences derived from the gene encoding the viral polyhedron protein, Friesen et al., (1986) "The Regulation of Baculovirus Gene Expression," in: *The Molecular Biology of Baculoviruses* (ed. Walter Doerfler); EPO Publ. Nos. 127 839 and 155 476; and the gene encoding the p10 protein, Vlak et al., (1988), *J. Gen. Virol.* 69:765.

DNA encoding suitable signal sequences can be derived from genes for secreted insect or baculovirus proteins, such as the baculovirus polyhedrin gene (Carbonell et al. (1988) *Gene*, 73:409). Alternatively, since the signals for mammalian cell posttranslational modifications (such as signal peptide cleavage, proteolytic cleavage, and phosphorylation) appear to be recognized by insect cells, and the signals required for secretion and nuclear accumulation also appear to be conserved between the invertebrate cells and vertebrate cells, leaders of non-insect origin, such as those derived from genes encoding human □-interferon, Maeda et al., (1985), *Nature 315*:592; human gastrin-releasing peptide, Lebacq-Verheyden et al., (1988), *Molec. Cell. Biol.* 8:3129; human IL-2, Smith et al., (1985) *Proc. Nat'l Acad. Sci. USA*, 82:8404;





mouse IL-3, (Miyajima et al., (1987) Gene 58:273; and human glucocerebrosidase, Martin et al. (1988) DNA, 7:99, can also be used to provide for secretion in insects.

A recombinant polypeptide or polyprotein may be expressed intracellularly or, if it is expressed with the proper regulatory sequences, it can be secreted. Good intracellular expression of nonfused foreign proteins usually requires heterologous genes that ideally have a short leader sequence containing suitable translation initiation signals preceding an ATG start signal. If desired, methionine at the N-terminus may be cleaved from the mature protein by *in vitro* incubation with cyanogen bromide.

Alternatively, recombinant polyproteins or proteins which are not naturally secreted can be secreted from the insect cell by creating chimeric DNA molecules that encode a fusion protein comprised of a leader sequence fragment that provides for secretion of the foreign protein in insects. The leader sequence fragment usually encodes a signal peptide comprised of hydrophobic amino acids which direct the translocation of the protein into the endoplasmic reticulum.

After insertion of the DNA sequence and/or the gene encoding the expression product precursor of the protein, an insect cell host is co-transformed with the heterologous DNA of the transfer vector and the genomic DNA of wild type baculovirus — usually by co-transfection. The promoter and transcription termination sequence of the construct will usually comprise a 2-5kb section of the baculovirus genome. Methods for introducing heterologous DNA into the desired site in the baculovirus virus are known in the art. (See Summers & Smith supra; Ju et al. (1987); Smith et al., Mol. Cell. Biol. (1983) 3:2156; and Luckow and Summers (1989)). For example, the insertion can be into a gene such as the polyhedrin gene, by homologous double crossover recombination; insertion can also be into a restriction enzyme site engineered into the desired baculovirus gene. Miller et al., (1989), Bioessays 4:91. The DNA sequence, when cloned in place of the polyhedrin gene in the expression vector, is flanked both 5' and 3' by polyhedrin-specific sequences and is positioned downstream of the polyhedrin promoter.

The newly formed baculovirus expression vector is subsequently packaged into an infectious recombinant baculovirus. Homologous recombination occurs at low frequency (between about 1% and about 5%); thus, the majority of the virus produced after cotransfection is still wild-type

virus. Therefore, a method is necessary to identify recombinant viruses. An advantage of the expression system is a visual screen allowing recombinant viruses to be distinguished. The polyhedrin protein, which is produced by the native virus, is produced at very high levels in the nuclei of infected cells at late times after viral infection. Accumulated polyhedrin protein forms occlusion bodies that also contain embedded particles. These occlusion bodies, up to 15 μm in size, are highly refractile, giving them a bright shiny appearance that is readily visualized under the light microscope. Cells infected with recombinant viruses lack occlusion bodies. To distinguish recombinant virus from wild-type virus, the transfection supernatant is plaqued onto a monolayer of insect cells by techniques known to those skilled in the art. Namely, the plaques are screened under the light microscope for the presence (indicative of wild-type virus) or absence (indicative of recombinant virus) of occlusion bodies. "Current Protocols in Microbiology" Vol. 2 (Ausubel et al. eds) at 16.8 (Supp. 10, 1990); Summers & Smith, *supra*; Miller et al. (1989).

Recombinant baculovirus expression vectors have been developed for infection into several insect cells. For example, recombinant baculoviruses have been developed for, inter alia: Aedes aegypti, Autographa californica, Bombyx mori, Drosophila melanogaster, Spodoptera frugiperda, and Trichoplusia ni (WO 89/046699; Carbonell et al., (1985) J. Virol. 56:153; Wright (1986) Nature 321:718; Smith et al., (1983) Mol. Cell. Biol. 3:2156; and see generally, Fraser, et al. (1989) In Vitro Cell. Dev. Biol. 25:225).

Cells and cell culture media are commercially available for both direct and fusion expression of heterologous polypeptides in a baculovirus/expression system; cell culture technology is generally known to those skilled in the art. See, eg. Summers & Smith supra.

The modified insect cells may then be grown in an appropriate nutrient medium, which allows for stable maintenance of the plasmid(s) present in the modified insect host. Where the expression product gene is under inducible control, the host may be grown to high density, and expression induced. Alternatively, where expression is constitutive, the product will be continuously expressed into the medium and the nutrient medium must be continuously circulated, while removing the product of interest and augmenting depleted nutrients. The product may be purified by such techniques as chromatography, eg. HPLC, affinity

chromatography, ion exchange chromatography, etc.; electrophoresis; density gradient centrifugation; solvent extraction, etc. As appropriate, the product may be further purified, as required, so as to remove substantially any insect proteins which are also present in the medium, so as to provide a product which is at least substantially free of host debris, eg. proteins, lipids and polysaccharides.

In order to obtain protein expression, recombinant host cells derived from the transformants are incubated under conditions which allow expression of the recombinant protein encoding sequence. These conditions will vary, dependent upon the host cell selected. However, the conditions are readily ascertainable to those of ordinary skill in the art, based upon what is known in the art.

#### iii. Plant Systems

There are many plant cell culture and whole plant genetic expression systems known in the art. Exemplary plant cellular genetic expression systems include those described in patents, such as: US 5,693,506; US 5,659,122; and US 5,608,143. Additional examples of genetic expression in plant cell culture has been described by Zenk, *Phytochemistry* 30:3861-3863 (1991).

Descriptions of plant protein signal peptides may be found in addition to the references described above in Vaulcombe et al., *Mol. Gen. Genet.* 209:33-40 (1987); Chandler et al., *Plant Molecular Biology* 3:407-418 (1984); Rogers, *J. Biol. Chem.* 260:3731-3738 (1985); Rothstein et al., *Gene* 55:353-356 (1987); Whittier et al., Nucleic Acids Research 15:2515-2535 (1987); Wirsel et al., *Molecular Microbiology* 3:3-14 (1989); Yu et al., *Gene* 122:247-253 (1992). A description of the regulation of plant gene expression by the phytohormone, gibberellic acid and secreted enzymes induced by gibberellic acid can be found in R.L. Jones and J. MacMillin, Gibberellins: in: *Advanced Plant Physiology*, Malcolm B. Wilkins, ed., 1984 Pitman Publishing Limited, London, pp. 21-52. References that describe other metabolically-regulated genes: Sheen, *Plant Cell*, 2:1027-1038(1990); Maas et al., *EMBO J.* 9:3447-3452 (1990); Benkel and Hickey, *Proc. Natl. Acad. Sci.* 84:1337-1339 (1987).

Typically, using techniques known in the art, a desired polynucleotide sequence is inserted into an expression cassette comprising genetic regulatory elements designed for operation in plants.

The expression cassette is inserted into a desired expression vector with companion sequences upstream and downstream from the expression cassette suitable for expression in a plant host. The companion sequences will be of plasmid or viral origin and provide necessary characteristics to the vector to permit the vectors to move DNA from an original cloning host, such as bacteria, to the desired plant host. The basic bacterial/plant vector construct will preferably provide a broad host range prokaryote replication origin; a prokaryote selectable marker; and, for Agrobacterium transformations, T DNA sequences for Agrobacterium-mediated transfer to plant chromosomes. Where the heterologous gene is not readily amenable to detection, the construct will preferably also have a selectable marker gene suitable for determining if a plant cell has been transformed. A general review of suitable markers, for example for the members of the grass family, is found in Wilmink and Dons, 1993, *Plant Mol. Biol. Reptr.*, 11(2):165-185.

Sequences suitable for permitting integration of the heterologous sequence into the plant genome are also recommended. These might include transposon sequences and the like for homologous recombination as well as Ti sequences which permit random insertion of a heterologous expression cassette into a plant genome. Suitable prokaryote selectable markers include resistance toward antibiotics such as ampicillin or tetracycline. Other DNA sequences encoding additional functions may also be present in the vector, as is known in the art. The nucleic acid molecules of the subject invention may be included into an expression cassette for expression of the protein(s) of interest. Usually, there will be only one expression cassette, although two or more are feasible. The recombinant expression cassette will contain in addition to the heterologous protein encoding sequence the following elements, a promoter region, plant 5' untranslated sequences, initiation codon depending upon whether or not the structural gene comes equipped with one, and a transcription and translation termination sequence. Unique restriction enzyme sites at the 5' and 3' ends of the cassette allow for easy insertion into a pre-existing vector.

A heterologous coding sequence may be for any protein relating to the present invention. The sequence encoding the protein of interest will encode a signal peptide which allows processing and translocation of the protein, as appropriate, and will usually lack any sequence which might

result in the binding of the desired protein of the invention to a membrane. Since, for the most part, the transcriptional initiation region will be for a gene which is expressed and translocated during germination, by employing the signal peptide which provides for translocation, one may also provide for translocation of the protein of interest. In this way, the protein(s) of interest will be translocated from the cells in which they are expressed and may be efficiently harvested. Typically secretion in seeds are across the aleurone or scutellar epithelium layer into the endosperm of the seed. While it is not required that the protein be secreted from the cells in which the protein is produced, this facilitates the isolation and purification of the recombinant protein.

Since the ultimate expression of the desired gene product will be in a eucaryotic cell it is desirable to determine whether any portion of the cloned gene contains sequences which will be processed out as introns by the host's splicosome machinery. If so, site-directed mutagenesis of the "intron" region may be conducted to prevent losing a portion of the genetic message as a false intron code, Reed and Maniatis, *Cell* 41:95-105, 1985.

The vector can be microinjected directly into plant cells by use of micropipettes to mechanically transfer the recombinant DNA. Crossway, *Mol. Gen. Genet*, 202:179-185, 1985. The genetic material may also be transferred into the plant cell by using polyethylene glycol, Krens, et al., *Nature*, 296, 72-74, 1982. Another method of introduction of nucleic acid segments is high velocity ballistic penetration by small particles with the nucleic acid either within the matrix of small beads or particles, or on the surface, Klein, et al., *Nature*, 327, 70-73, 1987 and Knudsen and Muller, 1991, *Planta*, 185:330-336 teaching particle bombardment of barley endosperm to create transgenic barley. Yet another method of introduction would be fusion of protoplasts with other entities, either minicells, cells, lysosomes or other fusible lipid-surfaced bodies, Fraley, et al., *Proc. Natl. Acad. Sci. USA*, 79, 1859-1863, 1982.

The vector may also be introduced into the plant cells by electroporation. (Fromm et al., *Proc. Natl Acad. Sci. USA* 82:5824, 1985). In this technique, plant protoplasts are electroporated in the presence of plasmids containing the gene construct. Electrical impulses of high field strength reversibly permeabilize biomembranes allowing the introduction of the plasmids. Electroporated plant protoplasts reform the cell wall, divide, and form plant callus.

All plants from which protoplasts can be isolated and cultured to give whole regenerated plants can be transformed by the present invention so that whole plants are recovered which contain the transferred gene. It is known that practically all plants can be regenerated from cultured cells or tissues, including but not limited to all major species of sugarcane, sugar beet, cotton, fruit and other trees, legumes and vegetables. Some suitable plants include, for example, species from the genera Fragaria, Lotus, Medicago, Onobrychis, Trifolium, Trigonella, Vigna, Citrus, Linum, Geranium, Manihot, Daucus, Arabidopsis, Brassica, Raphanus, Sinapis, Atropa, Capsicum, Datura, Hyoscyamus, Lycopersion, Nicotiana, Solanum, Petunia, Digitalis, Majorana, Cichorium, Helianthus, Lactuca, Bromus, Asparagus, Antirrhinum, Hererocallis, Nemesia, Pelargonium, Panicum, Pennisetum, Ranunculus, Senecio, Salpiglossis, Cucumis, Browaalia, Glycine, Lolium, Zea, Triticum, Sorghum, and Datura.

Means for regeneration vary from species to species of plants, but generally a suspension of transformed protoplasts containing copies of the heterologous gene is first provided. Callus tissue is formed and shoots may be induced from callus and subsequently rooted. Alternatively, embryo formation can be induced from the protoplast suspension. These embryos germinate as natural embryos to form plants. The culture media will generally contain various amino acids and hormones, such as auxin and cytokinins. It is also advantageous to add glutamic acid and proline to the medium, especially for such species as corn and alfalfa. Shoots and roots normally develop simultaneously. Efficient regeneration will depend on the medium, on the genotype, and on the history of the culture. If these three variables are controlled, then regeneration is fully reproducible and repeatable.

In some plant cell culture systems, the desired protein of the invention may be excreted or alternatively, the protein may be extracted from the whole plant. Where the desired protein of the invention is secreted into the medium, it may be collected. Alternatively, the embryos and embryoless-half seeds or other plant tissue may be mechanically disrupted to release any secreted protein between cells and tissues. The mixture may be suspended in a buffer solution to retrieve soluble proteins. Conventional protein isolation and purification methods will be then used to purify the recombinant protein. Parameters of time, temperature pH, oxygen, and

volumes will be adjusted through routine methods to optimize expression and recovery of heterologous protein.

#### iv. Bacterial Systems

Bacterial expression techniques are known in the art. A bacterial promoter is any DNA sequence capable of binding bacterial RNA polymerase and initiating the downstream (3') transcription of a coding sequence (eg. structural gene) into mRNA. A promoter will have a transcription initiation region which is usually placed proximal to the 5' end of the coding sequence. This transcription initiation region usually includes an RNA polymerase binding site and a transcription initiation site. A bacterial promoter may also have a second domain called an operator, that may overlap an adjacent RNA polymerase binding site at which RNA synthesis begins. The operator permits negative regulated (inducible) transcription, as a gene repressor protein may bind the operator and thereby inhibit transcription of a specific gene. Constitutive expression may occur in the absence of negative regulatory elements, such as the operator. In addition, positive regulation may be achieved by a gene activator protein binding sequence, which, if present is usually proximal (5') to the RNA polymerase binding sequence. An example of a gene activator protein is the catabolite activator protein (CAP), which helps initiate transcription of the lac operon in Escherichia coli (E. coli) [Raibaud et al. (1984) Annu. Rev. Genet. 18:173]. Regulated expression may therefore be either positive or negative, thereby either enhancing or reducing transcription.

Sequences encoding metabolic pathway enzymes provide particularly useful promoter sequences. Examples include promoter sequences derived from sugar metabolizing enzymes, such as galactose, lactose (lac) [Chang et al. (1977) Nature 198:1056], and maltose. Additional examples include promoter sequences derived from biosynthetic enzymes such as tryptophan (trp) [Goeddel et al. (1980) Nuc. Acids Res. 8:4057; Yelverton et al. (1981) Nucl. Acids Res. 9:731; US patent 4,738,921; EP-A-0036776 and EP-A-0121775]. The g-laotamase (bla) promoter system [Weissmann (1981) "The cloning of interferon and other mistakes." In Interferon 3 (ed. I. Gresser)], bacteriophage lambda PL [Shimatake et al. (1981) Nature

292:128] and T5 [US patent 4,689,406] promoter systems also provide useful promoter sequences.

In addition, synthetic promoters which do not occur in nature also function as bacterial promoters. For example, transcription activation sequences of one bacterial or bacteriophage promoter may be joined with the operon sequences of another bacterial or bacteriophage promoter, creating a synthetic hybrid promoter [US patent 4,551,433]. For example, the tac promoter is a hybrid trp-lac promoter comprised of both trp promoter and lac operon sequences that is regulated by the lac repressor [Amann et al. (1983) Gene 25:167; de Boer et al. (1983) Proc. Natl. Acad. Sci. 80:21]. Furthermore, a bacterial promoter can include naturally occurring promoters of non-bacterial origin that have the ability to bind bacterial RNA polymerase and initiate transcription. A naturally occurring promoter of non-bacterial origin can also be coupled with a compatible RNA polymerase to produce high levels of expression of some genes in prokaryotes. The bacteriophage T7 RNA polymerase/promoter system is an example of a coupled promoter system [Studier et al. (1986) J. Mol. Biol. 189:113; Tabor et al. (1985) Proc Natl. Acad. Sci. 82:1074]. In addition, a hybrid promoter can also be comprised of a bacteriophage promoter and an E. coli operator region (EPO-A-0 267 851). In addition to a functioning promoter sequence, an efficient ribosome binding site is also useful for the expression of foreign genes in prokaryotes. In E. coli, the ribosome binding site is called the Shine-Dalgarno (SD) sequence and includes an initiation codon (ATG) and a sequence 3-9 nucleotides in length located 3-11 nucleotides upstream of the initiation codon [Shine et al. (1975) Nature 254:34]. The SD sequence is thought to promote binding of mRNA to the ribosome by the pairing of bases between the SD sequence and the 3' and of E. coli 16S rRNA [Steitz et al. (1979) "Genetic signals and nucleotide sequences in messenger RNA." In Biological Regulation and Development: Gene Expression (ed. R.F. Goldberger)]. To express eukaryotic genes and prokaryotic genes with weak ribosome-binding site [Sambrook et al. (1989) "Expression of cloned genes in Escherichia coli." In Molecular Cloning: A Laboratory Manual].

A DNA molecule may be expressed intracellularly. A promoter sequence may be directly linked with the DNA molecule, in which case the first amino acid at the N-terminus will always

be a methionine, which is encoded by the ATG start codon. If desired, methionine at the N-terminus may be cleaved from the protein by *in vitro* incubation with cyanogen bromide or by either *in vivo* on *in vitro* incubation with a bacterial methionine N-terminal peptidase (EPO-A-0 219 237).

Fusion proteins provide an alternative to direct expression, Usually, a DNA sequence encoding the N-terminal portion of an endogenous bacterial protein, or other stable protein, is fused to the 5' end of heterologous coding sequences. Upon expression, this construct will provide a fusion of the two amino acid sequences. For example, the bacteriophage lambda cell gene can be linked at the 5' terminus of a foreign gene and expressed in bacteria. The resulting fusion protein preferably retains a site for a processing enzyme (factor Xa) to cleave the bacteriophage protein from the foreign gene [Nagai et al. (1984) Nature 309:810]. Fusion proteins can also be made with sequences from the lacZ [Jia et al. (1987) Gene 60:197], trpE [Allen et al. (1987) J. Biotechnol. 5:93; Makoff et al. (1989) J. Gen. Microbiol. 135:11], and Chey [EP-A-0 324 647] genes. The DNA sequence at the junction of the two amino acid sequences may or may not encode a cleavable site. Another example is a ubiquitin fusion protein. Such a fusion protein is made with the ubiquitin region that preferably retains a site for a processing enzyme (eg. ubiquitin specific processing-protease) to cleave the ubiquitin from the foreign protein. Through this method, native foreign protein can be isolated [Miller et al. (1989) Bio/Technology 7:698]. Alternatively, foreign proteins can also be secreted from the cell by creating chimeric DNA molecules that encode a fusion protein comprised of a signal peptide sequence fragment that provides for secretion of the foreign protein in bacteria [US patent 4,336,336]. The signal sequence fragment usually encodes a signal peptide comprised of hydrophobic amino acids which direct the secretion of the protein from the cell. The protein is either secreted into the growth media (gram-positive bacteria) or into the periplasmic space, located between the inner and outer membrane of the cell (gram-negative bacteria). Preferably there are processing sites, which can be cleaved either in vivo or in vitro encoded between the signal peptide fragment and the foreign gene.

DNA encoding suitable signal sequences can be derived from genes for secreted bacterial proteins, such as the *E. coli* outer membrane protein gene (*ompA*) [Masui *et al.* (1983), in:

Experimental Manipulation of Gene Expression; Ghrayeb et al. (1984) EMBO J. 3:2437] and the E. coli alkaline phosphatase signal sequence (phoA) [Oka et al. (1985) Proc. Natl. Acad. Sci. 82:7212]. As an additional example, the signal sequence of the alpha-amylase gene from various Bacillus strains can be used to secrete heterologous proteins from B. subtilis [Palva et al. (1982) Proc. Natl. Acad. Sci. USA 79:5582; EP-A-0 244 042].

Usually, transcription termination sequences recognized by bacteria are regulatory regions located 3' to the translation stop codon, and thus together with the promoter flank the coding sequence. These sequences direct the transcription of an mRNA which can be translated into the polypeptide encoded by the DNA. Transcription termination sequences frequently include DNA sequences of about 50 nucleotides capable of forming stem loop structures that aid in terminating transcription. Examples include transcription termination sequences derived from genes with strong promoters, such as the trp gene in E. coli as well as other biosynthetic genes. Usually, the above described components, comprising a promoter, signal sequence (if desired), coding sequence of interest, and transcription termination sequence, are put together into expression constructs. Expression constructs are often maintained in a replicon, such as an extrachromosomal element (eg. plasmids) capable of stable maintenance in a host, such as bacteria. The replicon will have a replication system, thus allowing it to be maintained in a prokaryotic host either for expression or for cloning and amplification. In addition, a replicon may be either a high or low copy number plasmid. A high copy number plasmid will generally have a copy number ranging from about 5 to about 200, and usually about 10 to about 150. A host containing a high copy number plasmid will preferably contain at least about 10, and more preferably at least about 20 plasmids. Either a high or low copy number vector may be selected, depending upon the effect of the vector and the foreign protein on the host.

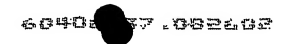
Alternatively, the expression constructs can be integrated into the bacterial genome with an integrating vector. Integrating vectors usually contain at least one sequence homologous to the bacterial chromosome that allows the vector to integrate. Integrations appear to result from recombinations between homologous DNA in the vector and the bacterial chromosome. For example, integrating vectors constructed with DNA from various Bacillus strains integrate into

the Bacillus chromosome (EP-A- 0 127 328). Integrating vectors may also be comprised of bacteriophage or transposon sequences.

Usually, extrachromosomal and integrating expression constructs may contain selectable markers to allow for the selection of bacterial strains that have been transformed. Selectable markers can be expressed in the bacterial host and may include genes which render bacteria resistant to drugs such as ampicillin, chloramphenicol, erythromycin, kanamycin (neomycin), and tetracycline [Davies et al. (1978) Annu. Rev. Microbiol. 32:469]. Selectable markers may also include biosynthetic genes, such as those in the histidine, tryptophan, and leucine biosynthetic pathways.

Alternatively, some of the above described components can be put together in transformation vectors. Transformation vectors are usually comprised of a selectable market that is either maintained in a replicon or developed into an integrating vector, as described above. Expression and transformation vectors, either extra-chromosomal replicons or integrating vectors, have been developed for transformation into many bacteria. For example, expression vectors have been developed for, inter alia, the following bacteria: Bacillus subtilis [Palva et al. (1982) Proc. Natl. Acad. Sci. USA 79:5582; EP-A-0 036 259 and EP-A-0 063 953; WO 84/04541], Escherichia coli [Shimatake et al. (1981) Nature 292:128; Amann et al. (1985) Gene 40:183; Studier et al. (1986) J. Mol. Biol. 189:113; EP-A-0 036 776,EP-A-0 136 829 and EP-A-0 136 907], Streptococcus cremoris [Powell et al. (1988) Appl. Environ. Microbiol. 54:655], Streptococcus lividans [Powell et al. (1988) Appl. Environ. Microbiol. 54:655], Streptomyces lividans [US patent 4,745,056].

Methods of introducing exogenous DNA into bacterial hosts are well-known in the art, and usually include either the transformation of bacteria treated with CaCl<sub>2</sub> or other agents, such as divalent cations and DMSO. DNA can also be introduced into bacterial cells by electroporation. Transformation procedures usually vary with the bacterial species to be transformed. See eg. [Masson et al. (1989) FEMS Microbiol. Lett. 60:273; Palva et al. (1982) Proc. Natl. Acad. Sci. USA 79:5582; EP-A-0 036 259 and EP-A-0 063 953; WO 84/04541, Bacillus], [Miller et al. (1988) Proc. Natl. Acad. Sci. 85:856; Wang et al. (1990) J. Bacteriol. 172:949, Campylobacter], [Cohen et al. (1973) Proc. Natl. Acad. Sci. 69:2110; Dower et al. (1988)



Nucleic Acids Res. 16:6127; Kushner (1978) "An improved method for transformation of Escherichia coli with ColE1-derived plasmids. In Genetic Engineering: Proceedings of the International Symposium on Genetic Engineering (eds. H.W. Boyer and S. Nicosia); Mandel et al. (1970) J. Mol. Biol. 53:159; Taketo (1988) Biochim. Biophys. Acta 949:318; Escherichia], [Chassy et al. (1987) FEMS Microbiol. Lett. 44:173 Lactobacillus]; [Fiedler et al. (1988) Anal. Biochem 170:38, Pseudomonas]; [Augustin et al. (1990) FEMS Microbiol. Lett. 66:203, Staphylococcus], [Barany et al. (1980) J. Bacteriol. 144:698; Harlander (1987) "Transformation of Streptococcus lactis by electroporation, in: Streptococcal Genetics (ed. J. Ferretti and R. Curtiss III); Perry et al. (1981) Infect. Immun. 32:1295; Powell et al. (1988) Appl. Environ. Microbiol. 54:655; Somkuti et al. (1987) Proc. 4th Evr. Cong. Biotechnology 1:412, Streptococcus].

## v. Yeast Expression

Yeast expression systems are also known to one of ordinary skill in the art. A yeast promoter is any DNA sequence capable of binding yeast RNA polymerase and initiating the downstream (3') transcription of a coding sequence (eg. structural gene) into mRNA. A promoter will have a transcription initiation region which is usually placed proximal to the 5' end of the coding sequence. This transcription initiation region usually includes an RNA polymerase binding site (the "TATA Box") and a transcription initiation site. A yeast promoter may also have a second domain called an upstream activator sequence (UAS), which, if present, is usually distal to the structural gene. The UAS permits regulated (inducible) expression. Constitutive expression occurs in the absence of a UAS. Regulated expression may be either positive or negative, thereby either enhancing or reducing transcription.

Yeast is a fermenting organism with an active metabolic pathway, therefore sequences encoding enzymes in the metabolic pathway provide particularly useful promoter sequences. Examples include alcohol dehydrogenase (ADH) (EP-A-0 284 044), enolase, glucokinase, glucose-6-phosphate isomerase, glyceraldehyde-3-phosphate-dehydrogenase (GAP or GAPDH), hexokinase, phosphofructokinase, 3-phosphoglycerate mutase, and pyruvate kinase (PyK)

(EPO-A-0 329 203). The yeast PHO5 gene, encoding acid phosphatase, also provides useful promoter sequences [Myanohara et al. (1983) Proc. Natl. Acad. Sci. USA 80:1]. In addition, synthetic promoters which do not occur in nature also function as yeast promoters. For example, UAS sequences of one yeast promoter may be joined with the transcription activation region of another yeast promoter, creating a synthetic hybrid promoter. Examples of such hybrid promoters include the ADH regulatory sequence linked to the GAP transcription activation region (US Patent Nos. 4,876,197 and 4,880,734). Other examples of hybrid promoters include promoters which consist of the regulatory sequences of either the ADH2, GAL4, GAL10, OR PHO5 genes, combined with the transcriptional activation region of a glycolytic enzyme gene such as GAP or PyK (EP-A-0 164 556). Furthermore, a yeast promoter can include naturally occurring promoters of non-yeast origin that have the ability to bind yeast RNA polymerase and initiate transcription. Examples of such promoters include, inter alia, [Cohen et al. (1980) Proc. Natl. Acad. Sci. USA 77:1078; Henikoff et al. (1981) Nature 283:835; Hollenberg et al. (1981) Curr. Topics Microbiol. Immunol. 96:119; Hollenberg et al. (1979) "The Expression of Bacterial Antibiotic Resistance Genes in the Yeast Saccharomyces cerevisiae," in: Plasmids of Medical, Environmental and Commercial Importance (eds. K.N. Timmis and A. Puhler); Mercerau-Puigalon et al. (1980) Gene 11:163; Panthier et al. (1980) Curr. Genet. 2:109;].

A DNA molecule may be expressed intracellularly in yeast. A promoter sequence may be directly linked with the DNA molecule, in which case the first amino acid at the N-terminus of the recombinant protein will always be a methionine, which is encoded by the ATG start codon. If desired, methionine at the N-terminus may be cleaved from the protein by *in vitro* incubation with cyanogen bromide.

Fusion proteins provide an alternative for yeast expression systems, as well as in mammalian, baculovirus, and bacterial expression systems. Usually, a DNA sequence encoding the N-terminal portion of an endogenous yeast protein, or other stable protein, is fused to the 5' end of heterologous coding sequences. Upon expression, this construct will provide a fusion of the two amino acid sequences. For example, the yeast or human superoxide dismutase (SOD) gene, can be linked at the 5' terminus of a foreign gene and expressed in yeast. The DNA sequence at the

junction of the two amino acid sequences may or may not encode a cleavable site. See eg. EP-A-0 196 056. Another example is a ubiquitin fusion protein. Such a fusion protein is made with the ubiquitin region that preferably retains a site for a processing enzyme (eg. ubiquitin-specific processing protease) to cleave the ubiquitin from the foreign protein. Through this method, therefore, native foreign protein can be isolated (eg. WO88/024066).

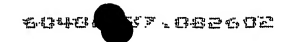
Alternatively, foreign proteins can also be secreted from the cell into the growth media by creating chimeric DNA molecules that encode a fusion protein comprised of a leader sequence fragment that provide for secretion in yeast of the foreign protein. Preferably, there are processing sites encoded between the leader fragment and the foreign gene that can be cleaved either *in vivo* or *in vitro*. The leader sequence fragment usually encodes a signal peptide comprised of hydrophobic amino acids which direct the secretion of the protein from the cell. DNA encoding suitable signal sequences can be derived from genes for secreted yeast proteins, such as the yeast invertase gene (EP-A-0 012 873; JPO. 62,096,086) and the A-factor gene (US patent 4,588,684). Alternatively, leaders of non-yeast origin, such as an interferon leader, exist that also provide for secretion in yeast (EP-A-0 060 057).

A preferred class of secretion leaders are those that employ a fragment of the yeast alpha-factor gene, which contains both a "pre" signal sequence, and a "pro" region. The types of alpha-factor fragments that can be employed include the full-length pre-pro alpha factor leader (about 83 amino acid residues) as well as truncated alpha-factor leaders (usually about 25 to about 50 amino acid residues) (US Patents 4,546,083 and 4,870,008; EP-A-0 324 274). Additional leaders employing an alpha-factor leader fragment that provides for secretion include hybrid alpha-factor leaders made with a presequence of a first yeast, but a pro-region from a second yeast alphafactor. (eg. see WO 89/02463.)

Usually, transcription termination sequences recognized by yeast are regulatory regions located 3' to the translation stop codon, and thus together with the promoter flank the coding sequence. These sequences direct the transcription of an mRNA which can be translated into the polypeptide encoded by the DNA. Examples of transcription terminator sequence and other yeast-recognized termination sequences, such as those coding for glycolytic enzymes.

Usually, the above described components, comprising a promoter, leader (if desired), coding sequence of interest, and transcription termination sequence, are put together into expression constructs. Expression constructs are often maintained in a replicon, such as an extrachromosomal element (eg. plasmids) capable of stable maintenance in a host, such as yeast or bacteria. The replicon may have two replication systems, thus allowing it to be maintained, for example, in yeast for expression and in a prokaryotic host for cloning and amplification. Examples of such yeast-bacteria shuttle vectors include YEp24 [Botstein et al. (1979) Gene 8:17-24], pCl/1 [Brake et al. (1984) Proc. Natl. Acad. Sci USA 81:4642-4646], and YRp17 [Stinchcomb et al. (1982) J. Mol. Biol. 158:157]. In addition, a replicon may be either a high or low copy number plasmid. A high copy number plasmid will generally have a copy number ranging from about 5 to about 200, and usually about 10 to about 150. A host containing a high copy number plasmid will preferably have at least about 10, and more preferably at least about 20. Enter a high or low copy number vector may be selected, depending upon the effect of the vector and the foreign protein on the host. See eg. Brake et al., supra.

Alternatively, the expression constructs can be integrated into the yeast genome with an integrating vector. Integrating vectors usually contain at least one sequence homologous to a yeast chromosome that allows the vector to integrate, and preferably contain two homologous sequences flanking the expression construct. Integrations appear to result from recombinations between homologous DNA in the vector and the yeast chromosome [Orr-Weaver et al. (1983) Methods in Enzymol. 101:228-245]. An integrating vector may be directed to a specific locus in yeast by selecting the appropriate homologous sequence for inclusion in the vector. See Orr-Weaver et al., supra. One or more expression construct may integrate, possibly affecting levels of recombinant protein produced [Rine et al. (1983) Proc. Natl. Acad. Sci. USA 80:6750]. The chromosomal sequences included in the vector can occur either as a single segment in the vector, which results in the integration of the entire vector, or two segments homologous to adjacent segments in the chromosome and flanking the expression construct in the vector, which can result in the stable integration of only the expression construct.



markers may include biosynthetic genes that can be expressed in the yeast host, such as ADE2, HIS4, LEU2, TRP1, and ALG7, and the G418 resistance gene, which confer resistance in yeast cells to tunicamycin and G418, respectively. In addition, a suitable selectable marker may also provide yeast with the ability to grow in the presence of toxic compounds, such as metal. For example, the presence of CUP1 allows yeast to grow in the presence of copper ions [Butt et al. (1987) Microbiol, Rev. 51:351].

Alternatively, some of the above described components can be put together into transformation vectors. Transformation vectors are usually comprised of a selectable marker that is either maintained in a replicon or developed into an integrating vector, as described above. Expression and transformation vectors, either extrachromosomal replicons or integrating vectors, have been developed for transformation into many yeasts. For example, expression vectors have been developed for, inter alia, the following yeasts: Candida albicans [Kurtz, et al. (1986) Mol. Cell. Biol. 6:142], Candida maltosa [Kunze, et al. (1985) J. Basic Microbiol. 25:141]. Hansenula polymorpha [Gleeson, et al. (1986) J. Gen. Microbiol. 132:3459; Roggenkamp et al. (1986) Mol. Gen. Genet. 202:302], Kluyveromyces fragilis [Das, et al. (1984) J. Bacteriol. 158:1165], Kluyveromyces lactis [De Louvencourt et al. (1983) J. Bacteriol. 154:737; Van den Berg et al. (1990) Bio/Technology 8:135], Pichia guillerimondii [Kunze et al. (1985) J. Basic Microbiol. 25:141], Pichia pastoris [Cregg, et al. (1985) Mol. Cell. Biol. 5:3376; US Patent Nos. 4,837,148 and 4,929,555], Saccharomyces cerevisiae [Hinnen et al. (1978) Proc. Natl. Acad. Sci. USA 75:1929; Ito et al. (1983) J. Bacteriol. 153:163], Schizosaccharomyces pombe [Beach and Nurse (1981) Nature 300:706], and Yarrowia lipolytica [Davidow, et al. (1985) Curr. Genet. 10:380471 Gaillardin, et al. (1985) Curr. Genet. 10:49].

Methods of introducing exogenous DNA into yeast hosts are well-known in the art, and usually include either the transformation of spheroplasts or of intact yeast cells treated with alkali cations. Transformation procedures usually vary with the yeast species to be transformed. See eg. [Kurtz et al. (1986) Mol. Cell. Biol. 6:142; Kunze et al. (1985) J. Basic Microbiol. 25:141; Candida]; [Gleeson et al. (1986) J. Gen. Microbiol. 132:3459; Roggenkamp et al. (1986) Mol. Gen. Genet. 202:302; Hansenula]; [Das et al. (1984) J. Bacteriol. 158:1165; De Louvencourt et

al. (1983) J. Bacteriol. 154:1165; Van den Berg et al. (1990) Bio/Technology 8:135;

Kluyveromyces]; [Cregg et al. (1985) Mol. Cell. Biol. 5:3376; Kunze et al. (1985) J. Basic

Microbiol. 25:141; US Patent Nos. 4,837,148 and 4,929,555; Pichia]; [Hinnen et al. (1978)

Proc. Natl. Acad. Sci. USA 75;1929; Ito et al. (1983) J. Bacteriol. 153:163 Saccharomyces];

[Beach and Nurse (1981) Nature 300:706; Schizosaccharomyces]; [Davidow et al. (1985) Curr.

Genet. 10:39; Gaillardin et al. (1985) Curr. Genet. 10:49; Yarrowia].

#### Antibodies

As used herein, the term "antibody" refers to a polypeptide or group of polypeptides composed of at least one antibody combining site. An "antibody combining site" is the three-dimensional binding space with an internal surface shape and charge distribution complementary to the features of an epitope of an antigen, which allows a binding of the antibody with the antigen. "Antibody" includes, for example, vertebrate antibodies, hybrid antibodies, chimeric antibodies, humanised antibodies, altered antibodies, univalent antibodies, Fab proteins, and single domain antibodies.

Antibodies against the proteins of the invention are useful for affinity chromatography, immunoassays, and distinguishing/identifying Streptococcal proteins.

Antibodies to the proteins of the invention, both polyclonal and monoclonal, may be prepared by conventional methods. In general, the protein is first used to immunize a suitable animal, preferably a mouse, rat, rabbit or goat. Rabbits and goats are preferred for the preparation of polyclonal sera due to the volume of serum obtainable, and the availability of labeled anti-rabbit and anti-goat antibodies. Immunization is generally performed by mixing or emulsifying the protein in saline, preferably in an adjuvant such as Freund's complete adjuvant, and injecting the mixture or emulsion parenterally (generally subcutaneously or intramuscularly). A dose of 50-200 µg/injection is typically sufficient. Immunization is generally boosted 2-6 weeks later with one or more injections of the protein in saline, preferably using Freund's incomplete adjuvant. One may alternatively generate antibodies by in vitro immunization using methods known in the art, which for the purposes of this invention is considered equivalent to in vivo immunization. Polyclonal antisera is obtained by bleeding the immunized animal into a glass or



plastic container, incubating the blood at 25  $\Box$  C for one hour, followed by incubating at 4  $\Box$  C for 2-18 hours. The serum is recovered by centrifugation (eg. 1,000g for 10 minutes). About 20-50 ml per bleed may be obtained from rabbits.

Monoclonal antibodies are prepared using the standard method of Kohler & Milstein [Nature (1975) 256:495-96], or a modification thereof. Typically, a mouse or rat is immunized as described above. However, rather than bleeding the animal to extract serum, the spleen (and optionally several large lymph nodes) is removed and dissociated into single cells. If desired, the spleen cells may be screened (after removal of nonspecifically adherent cells) by applying a cell suspension to a plate or well coated with the protein antigen. B-cells expressing membrane-bound immunoglobulin specific for the antigen bind to the plate, and are not rinsed away with the rest of the suspension. Resulting B-cells, or all dissociated spleen cells, are then induced to fuse with myeloma cells to form hybridomas, and are cultured in a selective medium (eg. hypoxanthine, aminopterin, thymidine medium, "HAT"). The resulting hybridomas are plated by limiting dilution, and are assayed for production of antibodies which bind specifically to the immunizing antigen (and which do not bind to unrelated antigens). The selected MAb-secreting hybridomas are then cultured either in vitro (eg. in tissue culture bottles or hollow fiber reactors), or in vivo (as ascites in mice).

If desired, the antibodies (whether polyclonal or monoclonal) may be labeled using conventional techniques. Suitable labels include fluorophores, chromophores, radioactive atoms (particularly <sup>32</sup>P and <sup>125</sup>I), electron-dense reagents, enzymes, and ligands having specific binding partners. Enzymes are typically detected by their activity. For example, horseradish peroxidase is usually detected by its ability to convert 3,3',5,5'-tetramethylbenzidine (TMB) to a blue pigment, quantifiable with a spectrophotometer. "Specific binding partner" refers to a protein capable of binding a ligand molecule with high specificity, as for example in the case of an antigen and a monoclonal antibody specific therefor. Other specific binding partners include biotin and avidin or streptavidin, IgG and protein A, and the numerous receptor-ligand couples known in the art. It should be understood that the above description is not meant to categorize the various labels into distinct classes, as the same label may serve in several different modes. For example, <sup>125</sup>I may serve as a radioactive label or as an electron-dense reagent. HRP may

serve as enzyme or as antigen for a MAb. Further, one may combine various labels for desired effect. For example, MAbs and avidin also require labels in the practice of this invention: thus, one might label a MAb with biotin, and detect its presence with avidin labeled with <sup>125</sup>I, or with an anti-biotin MAb labeled with HRP. Other permutations and possibilities will be readily apparent to those of ordinary skill in the art, and are considered as equivalents within the scope of the instant invention.

## Pharmaceutical Compositions

Pharmaceutical compositions can comprise either polypeptides, antibodies, or nucleic acid of the invention. The pharmaceutical compositions will comprise a therapeutically effective amount of either polypeptides, antibodies, or polynucleotides of the claimed invention. The term "therapeutically effective amount" as used herein refers to an amount of a therapeutic agent to treat, ameliorate, or prevent a desired disease or condition, or to exhibit a detectable therapeutic or preventative effect. The effect can be detected by, for example, chemical markers or antigen levels. Therapeutic effects also include reduction in physical symptoms, such as decreased body temperature. The precise effective amount for a subject will depend upon the subject's size and health, the nature and extent of the condition, and the therapeutics or combination of therapeutics selected for administration. Thus, it is not useful to specify an exact effective amount in advance. However, the effective amount for a given situation can be determined by routine experimentation and is within the judgement of the clinician.

For purposes of the present invention, an effective dose will be from about 0.01 mg/kg to 50 mg/kg or 0.05 mg/kg to about 10 mg/kg of the DNA constructs in the individual to which it is administered.

A pharmaceutical composition can also contain a pharmaceutically acceptable carrier. The term "pharmaceutically acceptable carrier" refers to a carrier for administration of a therapeutic agent, such as antibodies or a polypeptide, genes, and other therapeutic agents. The term refers to any pharmaceutical carrier that does not itself induce the production of antibodies harmful to the individual receiving the composition, and which may be administered without undue toxicity. Suitable carriers may be large, slowly metabolized macromolecules such as proteins,

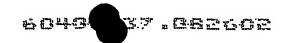
polysaccharides, polylactic acids, polyglycolic acids, polymeric amino acids, amino acid copolymers, and inactive virus particles. Such carriers are well known to those of ordinary skill in the art.

Pharmaceutically acceptable salts can be used therein, for example, mineral acid salts such as hydrochlorides, hydrobromides, phosphates, sulfates, and the like; and the salts of organic acids such as acetates, propionates, malonates, benzoates, and the like. A thorough discussion of pharmaceutically acceptable excipients is available in Remington's Pharmaceutical Sciences (Mack Pub. Co., N.J. 1991).

Pharmaceutically acceptable carriers in therapeutic compositions may contain liquids such as water, saline, glycerol and ethanol. Additionally, auxiliary substances, such as wetting or emulsifying agents, pH buffering substances, and the like, may be present in such vehicles. Typically, the therapeutic compositions are prepared as injectables, either as liquid solutions or suspensions; solid forms suitable for solution in, or suspension in, liquid vehicles prior to injection may also be prepared. Liposomes are included within the definition of a pharmaceutically acceptable carrier.

## **Delivery Methods**

Once formulated, the compositions of the invention can be administered directly to the subject. The subjects to be treated can be animals; in particular, human subjects can be treated. Direct delivery of the compositions will generally be accomplished by injection, either subcutaneously, intraperitoneally, intravenously or intramuscularly or delivered to the interstitial space of a tissue. The compositions can also be administered into a lesion. Other modes of administration include oral and pulmonary administration, suppositories, and transdermal or transcutaneous applications (eg. see WO98/20734), needles, and gene guns or hyposprays. Dosage treatment may be a single dose schedule or a multiple dose schedule. See also Delivery Strategies for Antisense Oligonucleotide Therapeutics (ed. Akhtar) ISBN 0849347785.



## **Vaccines**

Vaccines according to the invention may either be prophylactic (*ie.* to prevent infection) or therapeutic (*ie.* to treat disease after infection).

Such vaccines comprise immunising antigen(s), immunogen(s), polypeptide(s), protein(s) or nucleic acid, usually in combination with "pharmaceutically acceptable carriers," which include any carrier that does not itself induce the production of antibodies harmful to the individual receiving the composition. Suitable carriers are typically large, slowly metabolized macromolecules such as proteins, polysaccharides, polylactic acids, polyglycolic acids, polymeric amino acids, amino acid copolymers, lipid aggregates (such as oil droplets or liposomes), and inactive virus particles. Such carriers are well known to those of ordinary skill in the art. Additionally, these carriers may function as immunostimulating agents ("adjuvants"). Furthermore, the antigen or immunogen may be conjugated to a bacterial toxoid, such as a toxoid from diphtheria, tetanus, cholera, *H. pylori*, etc. pathogens.

Preferred adjuvants to enhance effectiveness of the composition include, but are not limited to: (1) aluminum salts (alum), such as aluminum hydroxide, aluminum phosphate, aluminum sulfate, etc; (2) oil-in-water emulsion formulations (with or without other specific immunostimulating agents such as muramyl peptides (see below) or bacterial cell wall components), such as for example (a) MF59™ (WO 90/14837; Chapter 10 in Vaccine design: the subunit and adjuvant approach, eds. Powell & Newman, Plenum Press 1995), containing 5% Squalene, 0.5% Tween 80, and 0.5% Span 85 (optionally containing various amounts of MTP-PE (see below), although not required) formulated into submicron particles using a microfluidizer such as Model 110Y microfluidizer (Microfluidics, Newton, MA), (b) SAF, containing 10% Squalane, 0.4% Tween 80, 5% pluronic-blocked polymer L121, and thr-MDP (see below) either microfluidized into a submicron emulsion or vortexed to generate a larger particle size emulsion, and (c) Ribi<sup>TM</sup> adjuvant system (RAS), (Ribi Immunochem, Hamilton, MT) containing 2% Squalene, 0.2% Tween 80, and one or more bacterial cell wall components from the group consisting of monophosphorylipid A (MPL), trehalose dimycolate (TDM), and cell wall skeleton (CWS), preferably MPL + CWS (Detox<sup>TM</sup>); (3) saponin adjuvants, such as Stimulon<sup>TM</sup> (Cambridge Bioscience, Worcester, MA) may be used or particles generated

therefrom such as ISCOMs (immunostimulating complexes); (4) Complete Freund's Adjuvant (CFA) and Incomplete Freund's Adjuvant (IFA); (5) cytokines, such as interleukins (eg. IL-1, IL-2, IL-4, IL-5, IL-6, IL-7, IL-12, etc.), interferons (eg. gamma interferon), macrophage colony stimulating factor (M-CSF), tumor necrosis factor (TNF), etc; and (6) other substances that act as immunostimulating agents to enhance the effectiveness of the composition. Alum and MF59<sup>TM</sup> are preferred.

As mentioned above, muramyl peptides include, but are not limited to, N-acetyl-muramyl-L-threonyl-D-isoglutamine (thr-MDP), N-acetyl-normuramyl-L-alanyl-D-isoglutamine (nor-MDP), N-acetylmuramyl-L-alanyl-D-isoglutaminyl-L-alanine-2-(1'-2'-dipalmitoyl-sn-glycero-3-hydroxyphosphoryloxy)-ethylamine (MTP-PE), etc.

The immunogenic compositions (eg. the immunising antigen/immunogen/polypeptide/protein/ nucleic acid, pharmaceutically acceptable carrier, and adjuvant) typically will contain diluents, such as water, saline, glycerol, ethanol, etc. Additionally, auxiliary substances, such as wetting or emulsifying agents, pH buffering substances, and the like, may be present in such vehicles. Typically, the immunogenic compositions are prepared as injectables, either as liquid solutions or suspensions; solid forms suitable for solution in, or suspension in, liquid vehicles prior to injection may also be prepared. The preparation also may be emulsified or encapsulated in liposomes for enhanced adjuvant effect, as discussed above under pharmaceutically acceptable carriers.

Immunogenic compositions used as vaccines comprise an immunologically effective amount of the antigenic or immunogenic polypeptides, as well as any other of the above-mentioned components, as needed. By "immunologically effective amount", it is meant that the administration of that amount to an individual, either in a single dose or as part of a series, is effective for treatment or prevention. This amount varies depending upon the health and physical condition of the individual to be treated, the taxonomic group of individual to be treated (eg. nonhuman primate, primate, etc.), the capacity of the individual's immune system to synthesize antibodies, the degree of protection desired, the formulation of the vaccine, the treating doctor's assessment of the medical situation, and other relevant factors. It is expected that the amount will fall in a relatively broad range that can be determined through routine trials.

The immunogenic compositions are conventionally administered parenterally, eg. by injection, either subcutaneously, intramuscularly, or transdermally/transcutaneously (eg. WO98/20734). Additional formulations suitable for other modes of administration include oral and pulmonary formulations, suppositories, and transdermal applications. Dosage treatment may be a single dose schedule or a multiple dose schedule. The vaccine may be administered in conjunction with other immunoregulatory agents.

As an alternative to protein-based vaccines, DNA vaccination may be used [eg. Robinson & Torres (1997) Seminars in Immunol 9:271-283; Donnelly et al. (1997) Annu Rev Immunol 15:617-648; later herein].

#### Gene Delivery Vehicles

Gene therapy vehicles for delivery of constructs including a coding sequence of a therapeutic of the invention, to be delivered to the mammal for expression in the mammal, can be administered either locally or systemically. These constructs can utilize viral or non-viral vector approaches in *in vivo* or *ex vivo* modality. Expression of such coding sequence can be induced using endogenous mammalian or heterologous promoters. Expression of the coding sequence in vivo can be either constitutive or regulated.

The invention includes gene delivery vehicles capable of expressing the contemplated nucleic acid sequences. The gene delivery vehicle is preferably a viral vector and, more preferably, a retroviral, adenoviral, adeno-associated viral (AAV), herpes viral, or alphavirus vector. The viral vector can also be an astrovirus, coronavirus, orthomyxovirus, papovavirus, paramyxovirus, picornavirus, poxvirus, or togavirus viral vector. See generally, Jolly (1994) Cancer Gene Therapy 1:51-64; Kimura (1994) Human Gene Therapy 5:845-852; Connelly (1995) Human Gene Therapy 6:185-193; and Kaplitt (1994) Nature Genetics 6:148-153.

Retroviral vectors are well known in the art and we contemplate that any retroviral gene therapy vector is employable in the invention, including B, C and D type retroviruses, xenotropic retroviruses (for example, NZB-X1, NZB-X2 and NZB9-1 (see O'Neill (1985) J. Virol. 53:160) polytropic retroviruses eg. MCF and MCF-MLV (see Kelly (1983) J. Virol. 45:291),

spumaviruses and lentiviruses. See RNA Tumor Viruses, Second Edition, Cold Spring Harbor Laboratory, 1985.

Portions of the retroviral gene therapy vector may be derived from different retroviruses. For example, retrovector LTRs may be derived from a Murine Sarcoma Virus, a tRNA binding site from a Rous Sarcoma Virus, a packaging signal from a Murine Leukemia Virus, and an origin of second strand synthesis from an Avian Leukosis Virus.

These recombinant retroviral vectors may be used to generate transduction competent retroviral vector particles by introducing them into appropriate packaging cell lines (see US patent 5,591,624). Retrovirus vectors can be constructed for site-specific integration into host cell DNA by incorporation of a chimeric integrase enzyme into the retroviral particle (see WO96/37626). It is preferable that the recombinant viral vector is a replication defective recombinant virus.

Packaging cell lines suitable for use with the above-described retrovirus vectors are well known in the art, are readily prepared (see WO95/30763 and WO92/05266), and can be used to create producer cell lines (also termed vector cell lines or "VCLs") for the production of recombinant vector particles. Preferably, the packaging cell lines are made from human parent cells (eg. HT1080 cells) or mink parent cell lines, which eliminates inactivation in human serum. Preferred retroviruses for the construction of retroviral gene therapy vectors include Avian Leukosis Virus, Bovine Leukemia, Virus, Murine Leukemia Virus, Mink-Cell Focus-Inducing Virus, Murine Sarcoma Virus, Reticuloendotheliosis Virus and Rous Sarcoma Virus. Particularly preferred Murine Leukemia Viruses include 4070A and 1504A (Hartley and Rowe (1976) J Virol 19:19-25), Abelson (ATCC No. VR-999), Friend (ATCC No. VR-245), Graffi, Gross (ATCC Nol VR-590), Kirsten, Harvey Sarcoma Virus and Rauscher (ATCC No. VR-998) and Moloney Murine Leukemia Virus (ATCC No. VR-190). Such retroviruses may be obtained from depositories or collections such as the American Type Culture Collection ("ATCC") in Rockville, Maryland or isolated from known sources using commonly available techniques.

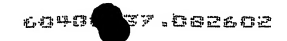
Exemplary known retroviral gene therapy vectors employable in this invention include those described in patent applications GB2200651, EP0415731, EP0345242, EP0334301,



WO89/02468; WO89/05349, WO89/09271, WO90/02806, WO90/07936, WO94/03622, WO93/25698, WO93/25234, WO93/11230, WO93/10218, WO91/02805, WO91/02825, WO95/07994, US 5,219,740, US 4,405,712, US 4,861,719, US 4,980,289, US 4,777,127, US 5,591,624. See also Vile (1993) Cancer Res 53:3860-3864; Vile (1993) Cancer Res 53:962-967; Ram (1993) Cancer Res 53 (1993) 83-88; Takamiya (1992) J Neurosci Res 33:493-503; Baba (1993) J Neurosurg 79:729-735; Mann (1983) Cell 33:153; Cane (1984) Proc Natl Acad Sci 81:6349; and Miller (1990) Human Gene Therapy 1. Human adenoviral gene therapy vectors are also known in the art and employable in this invention. See, for example, Berkner (1988) Biotechniques 6:616 and Rosenfeld (1991) Science 252:431, and WO93/07283, WO93/06223, and WO93/07282. Exemplary known adenoviral gene therapy vectors employable in this invention include those described in the above referenced documents and in WO94/12649, WO93/03769, WO93/19191, WO94/28938, WO95/11984, WO95/00655, WO95/27071, WO95/29993, WO95/34671, WO96/05320, WO94/08026, WO94/11506, WO93/06223, WO94/24299, WO95/14102, WO95/24297, WO95/02697, WO94/28152, WO94/24299, WO95/09241, WO95/25807, WO95/05835, WO94/18922 and WO95/09654. Alternatively, administration of DNA linked to killed adenovirus as described in Curiel (1992) Hum. Gene Ther. 3:147-154 may be employed. The gene delivery vehicles of the invention also include adenovirus associated virus (AAV) vectors. Leading and preferred examples of such vectors for use in this invention are the AAV-2 based vectors disclosed in Srivastava, WO93/09239. Most preferred AAV vectors comprise the two AAV inverted terminal repeats in which the native D-sequences are modified by substitution of nucleotides, such that at least 5 native nucleotides and up to 18 native nucleotides, preferably at least 10 native nucleotides up to 18 native nucleotides, most preferably 10 native nucleotides are retained and the remaining nucleotides of the D-sequence are deleted or replaced with non-native nucleotides. The native D-sequences of the AAV inverted terminal repeats are sequences of 20 consecutive nucleotides in each AAV inverted terminal repeat (ie. there is one sequence at each end) which are not involved in HP formation. The non-native replacement nucleotide may be any nucleotide other than the nucleotide found in the native D-sequence in the same position. Other employable exemplary AAV vectors are pWP-19, pWN-1, both of

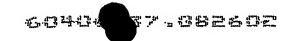
which are disclosed in Nahreini (1993) Gene 124:257-262. Another example of such an AAV vector is psub201 (see Samulski (1987) J. Virol. 61:3096). Another exemplary AAV vector is the Double-D ITR vector. Construction of the Double-D ITR vector is disclosed in US Patent 5,478,745. Still other vectors are those disclosed in Carter US Patent 4,797,368 and Muzyczka US Patent 5,139,941, Chartejee US Patent 5,474,935, and Kotin WO94/288157. Yet a further example of an AAV vector employable in this invention is SSV9AFABTKneo, which contains the AFP enhancer and albumin promoter and directs expression predominantly in the liver. Its structure and construction are disclosed in Su (1996) Human Gene Therapy 7:463-470. Additional AAV gene therapy vectors are described in US 5,354,678, US 5,173,414, US 5,139,941, and US 5,252,479.

The gene therapy vectors of the invention also include herpes vectors. Leading and preferred examples are herpes simplex virus vectors containing a sequence encoding a thymidine kinase polypeptide such as those disclosed in US 5,288,641 and EP0176170 (Roizman). Additional exemplary herpes simplex virus vectors include HFEM/ICP6-LacZ disclosed in WO95/04139 (Wistar Institute), pHSVlac described in Geller (1988) Science 241:1667-1669 and in WO90/09441 and WO92/07945, HSV Us3::pgC-lacZ described in Fink (1992) Human Gene Therapy 3:11-19 and HSV 7134, 2 RH 105 and GAL4 described in EP 0453242 (Breakefield), and those deposited with the ATCC with accession numbers VR-977 and VR-260. Also contemplated are alpha virus gene therapy vectors that can be employed in this invention. Preferred alpha virus vectors are Sindbis viruses vectors. Togaviruses, Semliki Forest virus (ATCC VR-67; ATCC VR-1247), Middleberg virus (ATCC VR-370), Ross River virus (ATCC VR-373; ATCC VR-1246), Venezuelan equine encephalitis virus (ATCC VR923; ATCC VR-1250; ATCC VR-1249; ATCC VR-532), and those described in US patents 5,091,309, 5,217,879, and WO92/10578. More particularly, those alpha virus vectors described in US Serial No. 08/405,627, filed March 15, 1995, WO94/21792, WO92/10578, WO95/07994, US 5,091,309 and US 5,217,879 are employable. Such alpha viruses may be obtained from depositories or collections such as the ATCC in Rockville, Maryland or isolated from known sources using commonly available techniques. Preferably, alphavirus vectors with reduced cytotoxicity are used (see USSN 08/679640).



DNA vector systems such as eukaryotic layered expression systems are also useful for expressing the nucleic acids of the invention. See WO95/07994 for a detailed description of eukaryotic layered expression systems. Preferably, the eukaryotic layered expression systems of the invention are derived from alphavirus vectors and most preferably from Sindbis viral vectors.

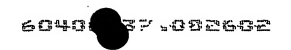
Other viral vectors suitable for use in the present invention include those derived from poliovirus, for example ATCC VR-58 and those described in Evans, Nature 339 (1989) 385 and Sabin (1973) J. Biol. Standardization 1:115; rhinovirus, for example ATCC VR-1110 and those described in Arnold (1990) J Cell Biochem L401; pox viruses such as canary pox virus or vaccinia virus, for example ATCC VR-111 and ATCC VR-2010 and those described in Fisher-Hoch (1989) Proc Natl Acad Sci 86:317; Flexner (1989) Ann NY Acad Sci 569:86, Flexner (1990) Vaccine 8:17; in US 4,603,112 and US 4,769,330 and WO89/01973; SV40 virus, for example ATCC VR-305 and those described in Mulligan (1979) Nature 277:108 and Madzak (1992) J Gen Virol 73:1533; influenza virus, for example ATCC VR-797 and recombinant influenza viruses made employing reverse genetics techniques as described in US 5,166,057 and in Enami (1990) Proc Natl Acad Sci 87:3802-3805; Enami & Palese (1991) J Virol 65:2711-2713 and Luytjes (1989) Cell 59:110, (see also McMichael (1983) NEJ Med 309:13, and Yap (1978) Nature 273:238 and Nature (1979) 277:108); human immunodeficiency virus as described in EP-0386882 and in Buchschacher (1992) J. Virol. 66:2731; measles virus, for example ATCC VR-67 and VR-1247 and those described in EP-0440219; Aura virus, for example ATCC VR-368; Bebaru virus, for example ATCC VR-600 and ATCC VR-1240; Cabassou virus, for example ATCC VR-922; Chikungunya virus, for example ATCC VR-64 and ATCC VR-1241; Fort Morgan Virus, for example ATCC VR-924; Getah virus, for example ATCC VR-369 and ATCC VR-1243; Kyzylagach virus, for example ATCC VR-927; Mayaro virus, for example ATCC VR-66; Mucambo virus, for example ATCC VR-580 and ATCC VR-1244; Ndumu virus, for example ATCC VR-371; Pixuna virus, for example ATCC VR-372 and ATCC VR-1245; Tonate virus, for example ATCC VR-925; Triniti virus, for example ATCC VR-469; Una virus, for example ATCC VR-374; Whataroa virus, for example ATCC VR-926; Y-62-33 virus, for example ATCC VR-375; O'Nyong virus,



Eastern encephalitis virus, for example ATCC VR-65 and ATCC VR-1242; Western encephalitis virus, for example ATCC VR-70, ATCC VR-1251, ATCC VR-622 and ATCC VR-1252; and coronavirus, for example ATCC VR-740 and those described in Hamre (1966) *Proc Soc Exp Biol Med* 121:190.

Delivery of the compositions of this invention into cells is not limited to the above mentioned viral vectors. Other delivery methods and media may be employed such as, for example, nucleic acid expression vectors, polycationic condensed DNA linked or unlinked to killed adenovirus alone, for example see US Serial No. 08/366,787, filed December 30, 1994 and Curiel (1992) Hum Gene Ther 3:147-154 ligand linked DNA, for example see Wu (1989) J Biol Chem 264:16985-16987, eucaryotic cell delivery vehicles cells, for example see US Serial No.08/240,030, filed May 9, 1994, and US Serial No. 08/404,796, deposition of photopolymerized hydrogel materials, hand-held gene transfer particle gun, as described in US Patent 5,149,655, ionizing radiation as described in US5,206,152 and in WO92/11033, nucleic charge neutralization or fusion with cell membranes. Additional approaches are described in Philip (1994) Mol Cell Biol 14:2411-2418 and in Woffendin (1994) Proc Natl Acad Sci 91:1581-1585.

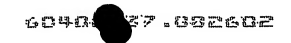
Particle mediated gene transfer may be employed, for example see US Serial No. 60/023,867. Briefly, the sequence can be inserted into conventional vectors that contain conventional control sequences for high level expression, and then incubated with synthetic gene transfer molecules such as polymeric DNA-binding cations like polylysine, protamine, and albumin, linked to cell targeting ligands such as asialoorosomucoid, as described in Wu & Wu (1987) *J. Biol. Chem.* 262:4429-4432, insulin as described in Hucked (1990) *Biochem Pharmacol* 40:253-263, galactose as described in Plank (1992) *Bioconjugate Chem* 3:533-539, lactose or transferrin. Naked DNA may also be employed. Exemplary naked DNA introduction methods are described in WO 90/11092 and US 5,580,859. Uptake efficiency may be improved using biodegradable latex beads. DNA coated latex beads are efficiently transported into cells after endocytosis initiation by the beads. The method may be improved further by treatment of the beads to increase hydrophobicity and thereby facilitate disruption of the endosome and release of the DNA into the cytoplasm.



Liposomes that can act as gene delivery vehicles are described in US 5,422,120, WO95/13796, WO94/23697, WO91/14445 and EP-524,968. As described in USSN. 60/023,867, on non-viral delivery, the nucleic acid sequences encoding a polypeptide can be inserted into conventional vectors that contain conventional control sequences for high level expression, and then be incubated with synthetic gene transfer molecules such as polymeric DNA-binding cations like polylysine, protamine, and albumin, linked to cell targeting ligands such as asialoorosomucoid, insulin, galactose, lactose, or transferrin. Other delivery systems include the use of liposomes to encapsulate DNA comprising the gene under the control of a variety of tissue-specific or ubiquitously-active promoters. Further non-viral delivery suitable for use includes mechanical delivery systems such as the approach described in Woffendin et al (1994) Proc. Natl. Acad. Sci. USA 91(24):11581-11585. Moreover, the coding sequence and the product of expression of such can be delivered through deposition of photopolymerized hydrogel materials. Other conventional methods for gene delivery that can be used for delivery of the coding sequence include, for example, use of hand-held gene transfer particle gun, as described in US 5,149,655; use of ionizing radiation for activating transferred gene, as described in US 5,206,152 and WO92/11033

Exemplary liposome and polycationic gene delivery vehicles are those described in US 5,422,120 and 4,762,915; in WO 95/13796; WO94/23697; and WO91/14445; in EP-0524968; and in Stryer, Biochemistry, pages 236-240 (1975) W.H. Freeman, San Francisco; Szoka (1980) Biochem Biophys Acta 600:1; Bayer (1979) Biochem Biophys Acta 550:464; Rivnay (1987) Meth Enzymol 149:119; Wang (1987) Proc Natl Acad Sci 84:7851; Plant (1989) Anal Biochem 176:420.

A polynucleotide composition can comprises therapeutically effective amount of a gene therapy vehicle, as the term is defined above. For purposes of the present invention, an effective dose will be from about 0.01 mg/kg to 50 mg/kg or 0.05 mg/kg to about 10 mg/kg of the DNA constructs in the individual to which it is administered.



## **Delivery Methods**

Once formulated, the polynucleotide compositions of the invention can be administered (1) directly to the subject; (2) delivered ex vivo, to cells derived from the subject; or (3) in vitro for expression of recombinant proteins. The subjects to be treated can be mammals or birds. Also, human subjects can be treated.

Direct delivery of the compositions will generally be accomplished by injection, either subcutaneously, intraperitoneally, intravenously or intramuscularly or delivered to the interstitial space of a tissue. The compositions can also be administered into a lesion. Other modes of administration include oral and pulmonary administration, suppositories, and transdermal or transcutaneous applications (eg. see WO98/20734), needles, and gene guns or hyposprays. Dosage treatment may be a single dose schedule or a multiple dose schedule. Methods for the ex vivo delivery and reimplantation of transformed cells into a subject are known in the art and described in eg. WO93/14778. Examples of cells useful in ex vivo applications include, for example, stem cells, particularly hematopoetic, lymph cells, macrophages, dendritic cells, or tumor cells.

Generally, delivery of nucleic acids for both ex vivo and in vitro applications can be accomplished by the following procedures, for example, dextran-mediated transfection, calcium phosphate precipitation, polybrene mediated transfection, protoplast fusion, electroporation, encapsulation of the polynucleotide(s) in liposomes, and direct microinjection of the DNA into nuclei, all well known in the art.

# Polynucleotide and polypeptide pharmaceutical compositions

The terms "polynucleotide" and "nucleic acid", used interchangeably herein,
In addition to the pharmaceutically acceptable carriers and salts described above, the following
additional agents can be used with polynucleotide and/or polypeptide compositions.

## A.Polypeptides

One example are polypeptides which include, without limitation: asioloorosomucoid (ASOR); transferrin; asialoglycoproteins; antibodies; antibody fragments; ferritin; interleukins;

interferons, granulocyte, macrophage colony stimulating factor (GM-CSF), granulocyte colony stimulating factor (G-CSF), macrophage colony stimulating factor (M-CSF), stem cell factor and erythropoietin. Viral antigens, such as envelope proteins, can also be used. Also, proteins from other invasive organisms, such as the 17 amino acid peptide from the circumsporozoite protein of plasmodium falciparum known as RII.

## B. Hormones, Vitamins, etc.

Other groups that can be included are, for example: hormones, steroids, androgens, estrogens, thyroid hormone, or vitamins, folic acid.

## C.Polyalkylenes, Polysaccharides, etc.

Also, polyalkylene glycol can be included with the desired polynucleotides/polypeptides. In a preferred embodiment, the polyalkylene glycol is polyethlylene glycol. In addition, mono-, di-, or polysaccharides can be included. In a preferred embodiment of this aspect, the polysaccharide is dextran or DEAE-dextran. Also, chitosan and poly(lactide-co-glycolide)

## D.Lipids, and Liposomes

The desired polynucleotide/polypeptide can also be encapsulated in lipids or packaged in liposomes prior to delivery to the subject or to cells derived therefrom.

Lipid encapsulation is generally accomplished using liposomes which are able to stably bind or entrap and retain nucleic acid. The ratio of condensed polynucleotide to lipid preparation can vary but will generally be around 1:1 (mg DNA:micromoles lipid), or more of lipid. For a review of the use of liposomes as carriers for delivery of nucleic acids, see, Hug and Sleight (1991) Biochim. Biophys. Acta. 1097:1-17; Straubinger (1983) Meth. Enzymol. 101:512-527. Liposomal preparations for use in the present invention include cationic (positively charged), anionic (negatively charged) and neutral preparations. Cationic liposomes have been shown to mediate intracellular delivery of plasmid DNA (Felgner (1987) Proc. Natl. Acad. Sci. USA 84:7413-7416); mRNA (Malone (1989) Proc. Natl. Acad. Sci. USA 86:6077-6081); and

purified transcription factors (Debs (1990) J. Biol. Chem. 265:10189-10192), in functional form.

Cationic liposomes are readily available. For example,

N[1-2,3-dioleyloxy)propyl]-N,N,N-triethylammonium (DOTMA) liposomes are available under the trademark Lipofectin, from GIBCO BRL, Grand Island, NY. (See, also, Felgner supra). Other commercially available liposomes include transfectace (DDAB/DOPE) and DOTAP/DOPE (Boerhinger). Other cationic liposomes can be prepared from readily available materials using techniques well known in the art. See, eg. Szoka (1978) Proc. Natl. Acad. Sci. USA 75:4194-4198; WO90/11092 for a description of the synthesis of DOTAP (1,2-bis(oleoyloxy)-3-(trimethylammonio)propane) liposomes.

Similarly, anionic and neutral liposomes are readily available, such as from Avanti Polar Lipids (Birmingham, AL), or can be easily prepared using readily available materials. Such materials include phosphatidyl choline, cholesterol, phosphatidyl ethanolamine, dioleoylphosphatidyl choline (DOPC), dioleoylphosphatidyl glycerol (DOPG), dioleoylphoshatidyl ethanolamine (DOPE), among others. These materials can also be mixed with the DOTMA and DOTAP starting materials in appropriate ratios. Methods for making liposomes using these materials are well known in the art.

The liposomes can comprise multilammelar vesicles (MLVs), small unilamellar vesicles (SUVs), or large unilamellar vesicles (LUVs). The various liposome-nucleic acid complexes are prepared using methods known in the art. See eg. Straubinger (1983) Meth. Immunol. 101:512-527; Szoka (1978) Proc. Natl. Acad. Sci. USA 75:4194-4198; Papahadjopoulos (1975) Biochim. Biophys. Acta 394:483; Wilson (1979) Cell 17:77); Deamer & Bangham (1976) Biochim. Biophys. Acta 443:629; Ostro (1977) Biochem. Biophys. Res. Commun. 76:836; Fraley (1979) Proc. Natl. Acad. Sci. USA 76:3348); Enoch & Strittmatter (1979) Proc. Natl. Acad. Sci. USA 76:145; Fraley (1980) J. Biol. Chem. (1980) 255:10431; Szoka & Papahadjopoulos (1978) Proc. Natl. Acad. Sci. USA 75:145; and Schaefer-Ridder (1982) Science 215:166.

#### **E.Lipoproteins**

In addition, lipoproteins can be included with the polynucleotide/polypeptide to be delivered. Examples of lipoproteins to be utilized include: chylomicrons, HDL, IDL, LDL, and VLDL. Mutants, fragments, or fusions of these proteins can also be used. Also, modifications of naturally occurring lipoproteins can be used, such as acetylated LDL. These lipoproteins can target the delivery of polynucleotides to cells expressing lipoprotein receptors. Preferably, if lipoproteins are including with the polynucleotide to be delivered, no other targeting ligand is included in the composition.

Naturally occurring lipoproteins comprise a lipid and a protein portion. The protein portion are known as apoproteins. At the present, apoproteins A, B, C, D, and E have been isolated and identified. At least two of these contain several proteins, designated by Roman numerals, AI, AII, AIV; CI, CII, CIII.

A lipoprotein can comprise more than one apoprotein. For example, naturally occurring chylomicrons comprises of A, B, C & E, over time these lipoproteins lose A and acquire C & E. VLDL comprises A, B, C & E apoproteins, LDL comprises apoprotein B; and HDL comprises apoproteins A, C, & E.

The amino acid of these apoproteins are known and are described in, for example, Breslow (1985) Annu Rev. Biochem 54:699; Law (1986) Adv. Exp Med. Biol. 151:162; Chen (1986) J Biol Chem 261:12918; Kane (1980) Proc Natl Acad Sci USA 77:2465; and Utermann (1984) Hum Genet 65:232.

Lipoproteins contain a variety of lipids including, triglycerides, cholesterol (free and esters), and phospholipids. The composition of the lipids varies in naturally occurring lipoproteins. For example, chylomicrons comprise mainly triglycerides. A more detailed description of the lipid content of naturally occurring lipoproteins can be found, for example, in *Meth. Enzymol.* 128 (1986). The composition of the lipids are chosen to aid in conformation of the apoprotein for receptor binding activity. The composition of lipids can also be chosen to facilitate hydrophobic interaction and association with the polynucleotide binding molecule.

Naturally occurring lipoproteins can be isolated from serum by ultracentrifugation, for instance. Such methods are described in *Meth. Enzymol.* (supra); Pitas (1980) J. Biochem.

255:5454-5460 and Mahey (1979) J Clin. Invest 64:743-750. Lipoproteins can also be produced by in vitro or recombinant methods by expression of the apoprotein genes in a desired host cell. See, for example, Atkinson (1986) Annu Rev Biophys Chem 15:403 and Radding (1958) Biochim Biophys Acta 30: 443. Lipoproteins can also be purchased from commercial suppliers, such as Biomedical Techniologies, Inc., Stoughton, Massachusetts, USA. Further description of lipoproteins can be found in Zuckermann et al. PCT/US97/14465.

## F.Polycationic Agents

Polycationic agents can be included, with or without lipoprotein, in a composition with the desired polynucleotide/polypeptide to be delivered.

Polycationic agents, typically, exhibit a net positive charge at physiological relevant pH and are capable of neutralizing the electrical charge of nucleic acids to facilitate delivery to a desired location. These agents have both in vitro, ex vivo, and in vivo applications. Polycationic agents can be used to deliver nucleic acids to a living subject either intramuscularly, subcutaneously, etc.

The following are examples of useful polypeptides as polycationic agents: polylysine, polyarginine, polyornithine, and protamine. Other examples include histones, protamines, human serum albumin, DNA binding proteins, non-histone chromosomal proteins, coat proteins from DNA viruses, such as (X174, transcriptional factors also contain domains that bind DNA and therefore may be useful as nucleic aid condensing agents. Briefly, transcriptional factors such as C/CEBP, c-jun, c-fos, AP-1, AP-2, AP-3, CPF, Prot-1, Sp-1, Oct-1, Oct-2, CREP, and TFIID contain basic domains that bind DNA sequences.

Organic polycationic agents include: spermine, spermidine, and purtrescine.

The dimensions and of the physical properties of a polycationic agent can be extrapolated from the list above, to construct other polypeptide polycationic agents or to produce synthetic polycationic agents.

Synthetic polycationic agents which are useful include, for example, DEAE-dextran, polybrene. Lipofectin, and lipofectAMINE are monomers that form polycationic complexes when combined with polynucleotides/polypeptides.



## Immunodiagnostic Assays

Streptococcus antigens of the invention can be used in immunoassays to detect antibody levels (or, conversely, anti-Streptococcus antibodies can be used to detect antigen levels). Immunoassays based on well defined, recombinant antigens can be developed to replace invasive diagnostics methods. Antibodies to Streptococcus proteins within biological samples, including for example, blood or serum samples, can be detected. Design of the immunoassays is subject to a great deal of variation, and a variety of these are known in the art. Protocols for the immunoassay may be based, for example, upon competition, or direct reaction, or sandwich type assays. Protocols may also, for example, use solid supports, or may be by immunoprecipitation. Most assays involve the use of labeled antibody or polypeptide; the labels may be, for example, fluorescent, chemiluminescent, radioactive, or dye molecules. Assays which amplify the signals from the probe are also known; examples of which are assays which utilize biotin and avidin, and enzyme-labeled and mediated immunoassays, such as ELISA assays.

Kits suitable for immunodiagnosis and containing the appropriate labeled reagents are constructed by packaging the appropriate materials, including the compositions of the invention, in suitable containers, along with the remaining reagents and materials (for example, suitable buffers, salt solutions, etc.) required for the conduct of the assay, as well as suitable set of assay instructions.

## Use of Polypeptides to Screen for Peptide Analogs and Antagonists

Polypeptides encoded by the instant polynucleotides and corresponding full length genes can be used to screen peptide libraries to identify binding partners, such as receptors, from within the library. Peptide libraries can be synthesized according to methods known in the art (e.g. Us patent 5,010,175; WO91/17823). Agonists or antagonists of the polypeptides if the invention can be screened using any available method known in the art, such as signal transduction, antibody binding, receptor binding, mitogenic assays, chemotaxis assays, etc. The assay conditions ideally should resemble the conditions under which the native activity is exhibited in vivo, that is, under physiologic pH, temperature, and ionic strength. Suitable agonists or

antagonists will exhibit strong inhibition or enhancement of the native activity at concentrations that do not cause toxic side effects in the subject. Agonists or antagonists that compete for binding to the native polypeptide can require concentrations equal to or greater than the native concentration, while inhibitors capable of binding irreversibly to the polypeptide can be added in concentrations on the order of the native concentration.

Such screening and experimentation can lead to identification of a polypeptide binding partner, such as a receptor, encoded by a gene or a cDNA corresponding to a polynucleotide described herein, and at least one peptide agonist or antagonist of the binding partner. Such agonists and antagonists can be used to modulate, enhance, or inhibit receptor function in cells to which the receptor is native, or in cells that possess the receptor as a result of genetic engineering. Further, if the receptor shares biologically important characteristics with a known receptor, information about agonist/antagonist binding can facilitate development of improved agonists/antagonists of the known receptor.

## Identification of anti-bacterial agents

# **Drug Screening Assays**

Of particular interest in the present invention is the identification of agents that have activity in modulating expression of one or more of the adhesion-specific genes described herein, so as to inhibit infection and/or disease. Of particular interest are screening assays for agents that have a low toxicity for human cells.

The term "agent" as used herein describes any molecule with the capability of altering or mimicking the expression or physiological function of a gene product of a differentially expressed gene. Generally a plurality of assay mixtures are run in parallel with different agent concentrations to obtain a differential response to the various concentrations. Typically, one of these concentrations serves as a negative control *i.e.* at zero concentration or below the level of detection.

Candidate agents encompass numerous chemical classes, including, but not limited to, organic molecules (e.g. small organic compounds having a molecular weight of more than 50 and less than about 2,500 daltons), peptides, antisense polynucleotides, and ribozymes, and the like.

Candidate agents can comprise functional groups necessary for structural interaction with proteins, particularly hydrogen bonding, and typically include at least an amine, carbonyl, hydroxyl or carboxyl group, preferably at least two of the functional chemical groups. The candidate agents often comprise cyclical carbon or heterocyclic structures and/or aromatic or polyaromatic structures substituted with one or more of the above functional groups. Candidate agents are also found among biomolecules including, but not limited to: polynucleotides, peptides, saccharides, fatty acids, steroids, purines, pyrimidines, derivatives, structural analogs or combinations thereof.

Candidate agents are obtained from a wide variety of sources including libraries of synthetic or natural compounds. For example, numerous means are available for random and directed synthesis of a wide variety of organic compounds and biomolecules, including expression of randomized oligonucleotides and oligopeptides. Alternatively, libraries of natural compounds in the form of bacterial, fungal, plant and animal extracts are available or readily produced. Additionally, natural or synthetically produced libraries and compounds are readily modified through conventional chemical, physical and biochemical means, and may be used to produce combinatorial libraries. Known pharmacological agents may be subjected to directed or random chemical modifications, such as acylation, alkylation, esterification, amidification, etc. to produce structural analogs.

## Screening of Candidate Agents In Vitro

A wide variety of *in vitro* assays may be used to screen candidate agents for the desired biological activity, including, but not limited to, labeled *in vitro* protein-protein binding assays, protein-DNA binding assays (e.g. to identify agents that affect expression), electrophoretic mobility shift assays, immunoassays for protein binding, and the like. For example, by providing for the production of large amounts of a differentially expressed polypeptide, one can identify ligands or substrates that bind to, modulate or mimic the action of the polypeptide. The purified polypeptide may also be used for determination of three-dimensional crystal structure, which can be used for modeling intermolecular interactions, transcriptional regulation, etc.

The screening assay can be a binding assay, wherein one or more of the molecules may be joined to a label, and the label directly or indirectly provide a detectable signal. Various labels include radioisotopes, fluorescers, chemiluminescers, enzymes, specific binding molecules, particles, e.g. magnetic particles, and the like. Specific binding molecules include pairs, such as biotin and streptavidin, digoxin and antidigoxin etc. For the specific binding members, the complementary member would normally be labeled with a molecule that provides for detection, in accordance with known procedures.

A variety of other reagents may be included in the screening assays described herein. Where the assay is a binding assay, these include reagents like salts, neutral proteins, e.g. albumin, detergents, etc. that are used to facilitate optimal protein-protein binding, protein-DNA binding, and/or reduce non-specific or background interactions. Reagents that improve the efficiency of the assay, such as protease inhibitors, nuclease inhibitors, anti-microbial agents, etc. may be used. The mixture of components are added in any order that provides for the requisite binding. Incubations are performed at any suitable temperature, typically between 4 and 40°C. Incubation periods are selected for optimum activity, but may also be optimized to facilitate rapid high-throughput screening. Typically between 0.1 and 1 hours will be sufficient. Many mammalian genes have homologs in yeast and lower animals. The study of such homologs' physiological role and interactions with other proteins in vivo or in vitro can facilitate understanding of biological function. In addition to model systems based on genetic complementation, yeast has been shown to be a powerful tool for studying protein-protein interactions through the two hybrid system.

#### Nucleic Acid Hybridisation

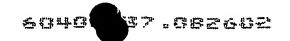
"Hybridization" refers to the association of two nucleic acid sequences to one another by hydrogen bonding. Typically, one sequence will be fixed to a solid support and the other will be free in solution. Then, the two sequences will be placed in contact with one another under conditions that favor hydrogen bonding. Factors that affect this bonding include: the type and volume of solvent; reaction temperature; time of hybridization; agitation; agents to block the non-specific attachment of the liquid phase sequence to the solid support (Denhardt's reagent or

BLOTTO); concentration of the sequences; use of compounds to increase the rate of association of sequences (dextran sulfate or polyethylene glycol); and the stringency of the washing conditions following hybridization. See Sambrook *et al.* [supra] Volume 2, chapter 9, pages 9.47 to 9.57.

"Stringency" refers to conditions in a hybridization reaction that favor association of very similar sequences over sequences that differ. For example, the combination of temperature and salt concentration should be chosen that is approximately 120 to 200 © below the calculated Tm of the hybrid under study. The temperature and salt conditions can often be determined empirically in preliminary experiments in which samples of genomic DNA immobilized on filters are hybridized to the sequence of interest and then washed under conditions of different stringencies. See Sambrook *et al.* at page 9.50.

Variables to consider when performing, for example, a Southern blot are (1) the complexity of the DNA being blotted and (2) the homology between the probe and the sequences being detected. The total amount of the fragment(s) to be studied can vary a magnitude of 10, from 0.1 to 1µg for a plasmid or phage digest to 10<sup>-9</sup> to 10<sup>-8</sup> g for a single copy gene in a highly' complex eukaryotic genome. For lower complexity polynucleotides, substantially shorter blotting, hybridization, and exposure times, a smaller amount of starting polynucleotides, and lower specific activity of probes can be used. For example, a single-copy yeast gene can be detected with an exposure time of only 1 hour starting with 1 µg of yeast DNA, blotting for two hours, and hybridizing for 4-8 hours with a probe of 10<sup>8</sup> cpm/µg. For a single-copy mammalian gene a conservative approach would start with 10 µg of DNA, blot overnight, and hybridize overnight in the presence of 10% dextran sulfate using a probe of greater than 10<sup>8</sup> cpm/µg, resulting in an exposure time of ~24 hours.

Several factors can affect the melting temperature (Tm) of a DNA-DNA hybrid between the probe and the fragment of interest, and consequently, the appropriate conditions for hybridization and washing. In many cases the probe is not 100% homologous to the fragment. Other commonly encountered variables include the length and total G+C content of the hybridizing sequences and the ionic strength and formamide content of the hybridization buffer. The effects of all of these factors can be approximated by a single equation:



Tm= 81 + 16.6(log<sub>10</sub>Ci) + 0.4[%(G + C)]-0.6(%formamide) - 600/n-1.5(%mismatch). where Ci is the salt concentration (monovalent ions) and n is the length of the hybrid in base pairs (slightly modified from Meinkoth & Wahl (1984) Anal. Biochem. 138: 267-284). In designing a hybridization experiment, some factors affecting nucleic acid hybridization can be conveniently altered. The temperature of the hybridization and washes and the salt concentration during the washes are the simplest to adjust. As the temperature of the hybridization increases (ie. stringency), it becomes less likely for hybridization to occur between strands that are nonhomologous, and as a result, background decreases. If the radiolabeled probe is not completely homologous with the immobilized fragment (as is frequently the case in gene family and interspecies hybridization experiments), the hybridization temperature must be reduced, and background will increase. The temperature of the washes affects the intensity of the hybridizing band and the degree of background in a similar manner. The stringency of the washes is also increased with decreasing salt concentrations.

In general, convenient hybridization temperatures in the presence of 50% formamide are 42°C for a probe with is 95% to 100% homologous to the target fragment, 37°C for 90% to 95% homology, and 32°C for 85% to 90% homology. For lower homologies, formamide content should be lowered and temperature adjusted accordingly, using the equation above. If the homology between the probe and the target fragment are not known, the simplest approach is to start with both hybridization and wash conditions which are nonstringent. If non-specific bands or high background are observed after autoradiography, the filter can be washed at high stringency and reexposed. If the time required for exposure makes this approach impractical, several hybridization and/or washing stringencies should be tested in parallel.

# Nucleic Acid Probe Assays

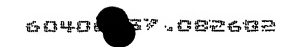
Methods such as PCR, branched DNA probe assays, or blotting techniques utilizing nucleic acid probes according to the invention can determine the presence of cDNA or mRNA. A probe is said to "hybridize" with a sequence of the invention if it can form a duplex or double stranded complex, which is stable enough to be detected.



The nucleic acid probes will hybridize to the Streptococcus nucleotide sequences of the invention (including both sense and antisense strands). Though many different nucleotide sequences will encode the amino acid sequence, the native Streptococcal sequence is preferred because it is the actual sequence present in cells. mRNA represents a coding sequence and so a probe should be complementary to the coding sequence; single-stranded cDNA is complementary to mRNA, and so a cDNA probe should be complementary to the non-coding sequence.

The probe sequence need not be identical to the Streptococcal sequence (or its complement) some variation in the sequence and length can lead to increased assay sensitivity if the nucleic acid probe can form a duplex with target nucleotides, which can be detected. Also, the nucleic acid probe can include additional nucleotides to stabilize the formed duplex. Additional Streptococcus sequence may also be helpful as a label to detect the formed duplex. For example, a non-complementary nucleotide sequence may be attached to the 5' end of the probe. with the remainder of the probe sequence being complementary to a Streptococcus sequence. Alternatively, non-complementary bases or longer sequences can be interspersed into the probe, provided that the probe sequence has sufficient complementarity with the a Streptococcus sequence in order to hybridize therewith and thereby form a duplex which can be detected. The exact length and sequence of the probe will depend on the hybridization conditions (e.g. temperature, salt condition etc.). For example, for diagnostic applications, depending on the complexity of the analyte sequence, the nucleic acid probe typically contains at least 10-20 nucleotides, preferably 15-25, and more preferably at least 30 nucleotides, although it may be shorter than this. Short primers generally require cooler temperatures to form sufficiently stable hybrid complexes with the template.

Probes may be produced by synthetic procedures, such as the triester method of Matteucci et al. [J. Am. Chem. Soc. (1981) 103:3185], or according to Urdea et al. [Proc. Natl. Acad. Sci. USA (1983) 80: 7461], or using commercially available automated oligonucleotide synthesizers. The chemical nature of the probe can be selected according to preference. For certain applications, DNA or RNA are appropriate. For other applications, modifications may be incorporated eg. backbone modifications, such as phosphorothioates or methylphosphonates,



can be used to increase in vivo half-life, alter RNA affinity, increase nuclease resistance etc. [eg. see Agrawal & Iyer (1995) Curr Opin Biotechnol 6:12-19; Agrawal (1996) TIBTECH 14:376-387]; analogues such as peptide nucleic acids may also be used [eg. see Corey (1997) TIBTECH 15:224-229; Buchardt et al. (1993) TIBTECH 11:384-386].

Alternatively, the polymerase chain reaction (PCR) is another well-known means for detecting small amounts of target nucleic acid. The assay is described in Mullis *et al.* [*Meth. Enzymol.* (1987) 155:335-350] & US patents 4,683,195 & 4,683,202. Two "primer" nucleotides hybridize with the target nucleic acids and are used to prime the reaction. The primers can comprise sequence that does not hybridize to the sequence of the amplification target (or its complement) to aid with duplex stability or, for example, to incorporate a convenient restriction site. Typically, such sequence will flank the desired Streptococcus sequence.

A thermostable polymerase creates copies of target nucleic acids from the primers using the original target nucleic acids as a template. After a threshold amount of target nucleic acids are generated by the polymerase, they can be detected by more traditional methods, such as Southern blots. When using the Southern blot method, the labelled probe will hybridize to the Streptococcus sequence (or its complement).

Also, mRNA or cDNA can be detected by traditional blotting techniques described in Sambrook et al [supra]. mRNA, or cDNA generated from mRNA using a polymerase enzyme, can be purified and separated using gel electrophoresis. The nucleic acids on the gel are then blotted onto a solid support, such as nitrocellulose. The solid support is exposed to a labelled probe and then washed to remove any unhybridized probe. Next, the duplexes containing the labelled probe are detected. Typically, the probe is labelled with a radioactive moiety.

Figure 1

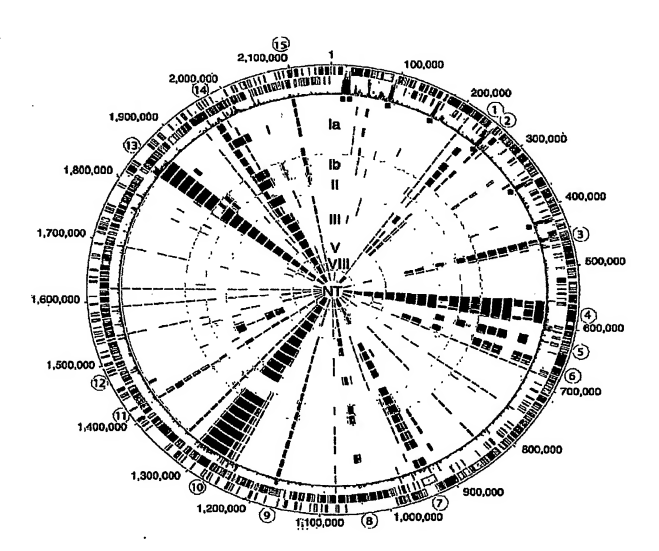


Figure 2

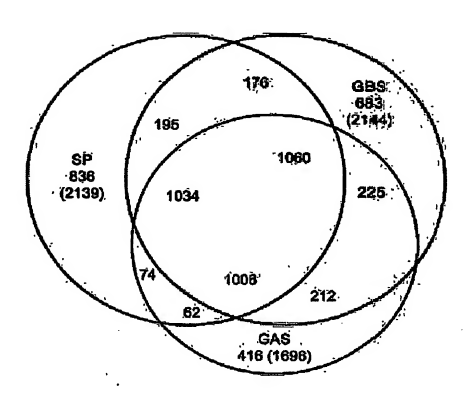
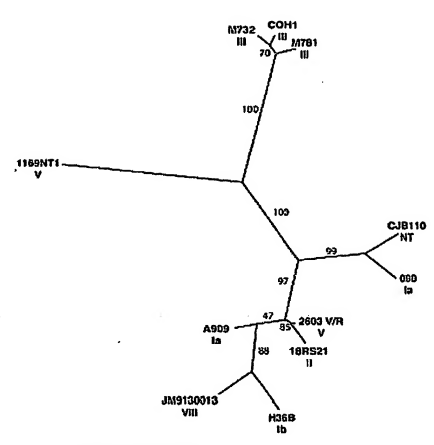


Figure 3

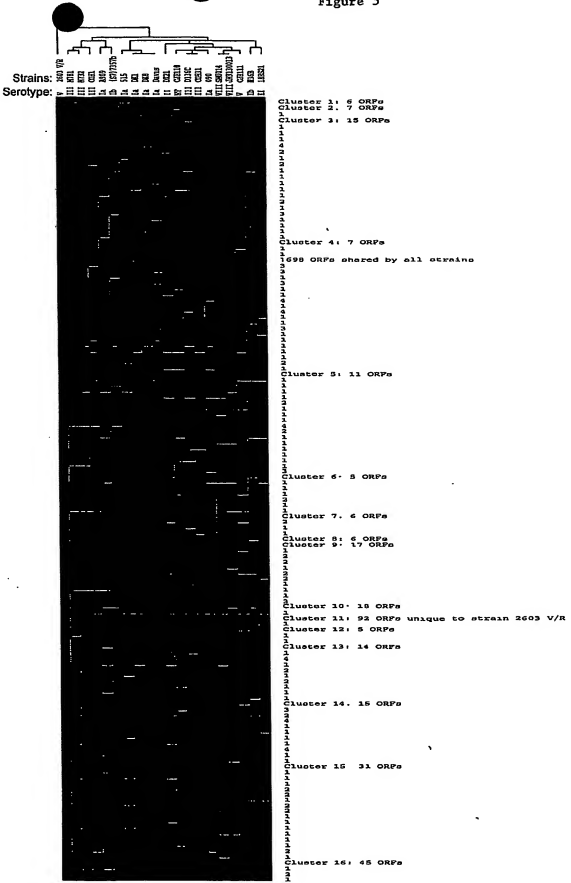


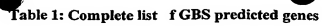
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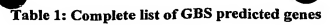




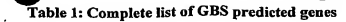
ORF	Size	Ann tati n	
	(a.a.)		
SAG0001	453	chromosomal replication initiator protein DnaA	
SAG0002	378	DNA polymerase III, beta subunit	
SAG0003	293	diacylglycerol kinase catalytic domain protein, putative	
SAG0004	65	conserved hypothetical protein	
SAG0005	67	hypothetical protein	
SAG0006	371	GTP-binding protein YchF	
SAG0007	191	peptidyl-tRNA hydrolase	
SAG0008	1165	transcription-repair coupling factor	
SAG0009	31	hypothetical protein	
SAG0010	90	S4 domain protein	
SAG0011	123	cell division protein DivIC, putative	
SAG0012	44	conserved hypothetical protein	
SAG0013	428	protein of unknown function	
SAG0014	424	MesJ/Ycf62 family protein	
SAG0015	180	hypoxanthine-guanine phosphoribosyltransferase	
SAG0016	658	cell division protein FtsH	
SAG0017	447	pcsB protein	
SAG0017	322	ribose-phosphate pyrophosphokinase	
SAG0019	391	aminotransferase, class I	
SAG0019	253	recombination protein O	
SAG0020 SAG0021	283		
SAG0021	330	protease, putative	
SAG0022 SAG0023		fatty acid/phospholipid synthesis protein PlsX	
	79 234	acyl carrier protein	
SAG0024		phosphoribosylaminoimidazole-succinocarboxamide synthase	
SAG0025	1241	phosphoribosylformylglycinamidine synthase, putative	
SAG0026	484	amidophosphoribosyltransferase	
SAG0027	340	phosphoribosylformylglycinamidine cyclo-ligase	
SAG0028	182	phosphoribosylglycinamide formyltransferase	
SAG0029	250	acetyltransferase, GNAT family	
SAG0030	515	phosphoribosylaminoimidazolecarboxamide	
0.4.00021	200	formyltransferase/IMP cyclohydrolase	
SAG0031	299	peptidase, M23/M37 family	
SAG0032	434	group B streptococcal surface immunogenic protein	
SAG0033	232	N-acetylmannosamine-6-P epimerase, putative	
SAG0034	438	sugar ABC transporter, sugar-binding protein	
SAG0035	295		
SAG0036	276	<u> </u>	
SAG0037	147	conserved hypothetical protein	
SAG0038	220		
SAG0039	305		
SAG0040	293		
SAG0041	325		
SAG0042	267	phosphosugar-binding transcriptional regulator, RpiR family,	
		putative	
SAG0043	421	phosphoribosylamineglycine ligase	
SAG0044	162	phosphoribosylaminoimidazole carboxylase, catalytic subunit	
SAG0045	363	phosphoribosylaminoimidazole carboxylase, ATPase subunit	
SAG0046	463	membrane protein, putative	



ORF	Size	Annotation
	(a.a.)	
SAG0047	432	adenylosuccinate lyase
SAG0048	303	transcriptional regulator, Cro/CI family
SAG0049	332	Holliday junction DNA helicase RuvB
SAG0050	145	phosphotyrosine protein phosphatase, low molecular weight
SAG0051	126	MORN motif family protein
SAG0052	592	membrane protein, putative
SAG0053	880	aldehyde-alcohol dehydrogenase
SAG0054	338	alcohol dehydrogenase, propanol-preferring
SAG0055	496	threonine synthase
SAG0056	412	MATE efflux family protein
SAG0057	102	ribosomal protein S10
SAG0058	208	ribosomal protein L3
SAG0059	207	ribosomal protein L4
SAG0060	98	ribosomal protein L23
SAG0061	277	ribosomal protein L2
SAG0062	92	ribosomal protein S19
. SAG0063	114	ribosomal protein L22
. SAG0064	217	ribosomal protein S3
SAG0065	137	ribosomal protein L16
SAG0066	68	ribosomal protein L29
SAG0067	86	ribosomal protein S17
SAG0068	122	ribosomal protein L14
SAG0069	101	ribosomal protein L24
SAG0070	180	ribosomal protein L5
SAG0071	61	ribosomal protein S14, putative
SAG0072	132	ribosomal protein S8
SAG0073	178	ribosomal protein L6
SAG0074	118	ribosomal protein L18
SAG0075	164	ribosomal protein S5
SAG0076	59	ribosomal protein L30
SAG0077	146	ribosomal protein L15
SAG0078	434	preprotein translocase, SecY subunit
SAG0079	212	adenylate kinase
SAG0080	72	translation initiation factor IF-1
SAG0081	38	<u> </u>
SAG0082	121	ribosomal protein S13
SAG0083	118	
SAG0084	312	
SAG0085	128	
SAG0086	85	<u> </u>
SAG0087	59	
SAG0088	56	<u> </u>
SAG0089	183	conserved hypothetical protein
SAG0090	139	conserved hypothetical protein
SAG0091	144	· · · · · · · · · · · · · · · · · · ·
SAG0092	230	
SAG0093	250	
SAG0094	191	N-acetylmuramoyl-L-alanine amidase, family 4 protein



ORF	Size	Annotation
	(a.a.)	
SAG0095	344	heat-inducible transcription repressor HrcA
SAG0096	190	heat shock protein GrpE
SAG0097	609	dnaK protein
SAG0098	379	dnaJ protein
SAG0099	415	transcriptional regulator, GntR family
SAG0100	258	tRNA pseudouridine synthase A .
SAG0101	252	phosphomethylpyrimidine kinase, putative
SAG0102	154	conserved hypothetical protein
SAG0103	189	conserved hypothetical protein TIGR01440
SAG0104	280	conserved hypothetical protein
SAG0105	427	trigger factor
SAG0106	191	DNA-directed RNA polymerase, delta subunit, putative
SAG0107	534	CTP synthase
SAG0108	308	conserved hypothetical protein
SAG0109	148	deoxyuridine 5'-triphosphate nucleotidohydrolase
SAG0110	454	DNA repair protein RadA
SAG0111	165	carbonic anhydrase-related protein
SAG0112	439	pyridine nucleotide-disulphide oxidoreductase family protein
SAG0113	484	glutamyl-tRNA synthetase
SAG0114	322	ribose ABC transporter, periplasmic D-ribose-binding protein
SAG0115	310	ribose ABC transporter, permease protein
SAG0116	492	ribose ABC transporter, ATP-binding protein
SAG0117	132	ribose ABC transporter protein RbsD
SAG0118	303	ribokinase
SAG0119	328	ribose operon repressor RbsR
SAG0120	32	hypothetical protein
SAG0121	362	permease, putative
SAG0122	228	
SAG0123	223	
SAG0124	356	
SAG0125	396	
SAG0126	462	argininosuccinate lyase
SAG0127	293	fructose-bisphosphate aldolase
SAG0128	305	
SAG0129	62	
SAG0130	121	conserved hypothetical protein
SAG0131	543	
SAG0132	294	
SAG0132	38	
SAG0134	96	<del> </del>
SAG0134	246	
SAG0136	516	
51130130	510	protein
SAG0137	627	- <del>                                    </del>
SAG0137	279	
SAG0138	251	
SAG0139	386	
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ORF	Size	Annotation
	(a.a.)	
SAG0142	420	conserved hypothetical protein
SAG0143	410	selenocysteine lyase
SAG0144	147	NifU family protein
SAG0145	472	conserved hypothetical protein
SAG0146	395	penicillin-binding protein 4, putative
SAG0147	411	D-alanyl-D-alanine carboxypeptidase family protein
SAG0148	551	oligopeptide ABC transporter, substrate-binding protein, putative
SAG0149	304	oligopeptide ABC transporter, permease protein
SAG0150	343	oligopeptide ABC transporter, permease protein
SAG0151	348	oligopeptide ABC transporter, ATP-binding protein
SAG0152	310	oligopeptide ABC transporter, ATP-binding protein
SAG0153	283	4-diphosphocytidyl-2C-methyl-D-erythritol kinase
SAG0154	147	adc operon repressor AdcR
SAG0155	236	zinc ABC transporter, ATP-binding protein
SAG0156	270	zinc ABC transporter, permease protein
SAG0157	NA	deoxyribonuclease-related protein, degenerate
SAG0158	419	tyrosyl-tRNA synthetase
SAG0159	765	penicillin-binding protein 1B, putative
SAG0160	1191	DNA-directed RNA polymerase, beta subunit
SAG0161	1216	DNA-directed RNA polymerase beta' subunit
SAG0162	121	conserved hypothetical protein
SAG0163	323	competence protein CglA
SAG0164	282	competence protein CglB
SAG0165	151	conserved hypothetical protein
SAG0166	123	conserved domain protein
SAG0167	324	conserved hypothetical protein
SAG0168	397	acetate kinase
SAG0169	68	transcriptional regulator, Cro/CI family
SAG0170	45	hypothetical protein
SAG0171	151	hypothetical protein
SAG0172	221	protease, putative
SAG0173	256	pyrroline-5-carboxylate reductase
SAG0174	355	glutamyl-aminopeptidase
SAG0175	79	hypothetical protein
SAG0176	94	conserved hypothetical protein
SAG0177	107	thioredoxin family protein
SAG0178	208	tRNA binding domain protein
SAG0179	·238	conserved hypothetical protein
SAG0180	131	
SAG0181	214	hydrolase, haloacid dehalogenase-like family
SAG0182	581	
SAG0183	246	
SAG0184	151	
SAG0185	242	
SAG0186	36	
SAG0187	542	
SAG0188	325	
SAG0189		

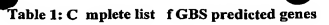


ORF	Size	Annotati n
	(a.a.)	
SAG0190	267	peptide ABC transporter, ATP-binding protein
SAG0191	208	peptide ABC transporter, ATP-binding protein
SAG0192	676	PTS system, IIABC components
SAG0193	541	alpha amylase family protein
SAG0194	639	transcriptional antiterminator, BglG family
SAG0195	377	IS1548, transposase
SAG0196	66	conserved domain protein
SAG0197	94	PTS system, IIB component, putative
SAG0198	451	PTS system, IIC component, putative
SAG0199	285	transketolase, N-terminal subunit
SAG0200	309	transketolase, C-terminal subunit
SAG0201	419	oxidoreductase, putative
SAG0202	89	ribosomal protein S15
SAG0203	709	polyribonucleotide nucleotidyltransferase
SAG0204	250	conserved hypothetical protein
SAG0205	194	serine O-acetyltransferase
SAG0206	60	lipoprotein, putative
SAG0207	447	cysteinyl-tRNA synthetase
SAG0208	128	conserved hypothetical protein
SAG0209	251	RNA methyltransferase, TrmH family, group 3
SAG0210	172	conserved hypothetical protein
SAG0210	286	DegV family protein
SAG0211	32	hypothetical protein
SAG0213	39	
SAG0213	148	ribosomal protein L13
SAG0214	130	ribosomal protein S9
SAG0216	33	hypothetical protein
SAG0217	384	
SAG0217	158	transcriptional regulator, Cro/CI family
SAG0219	101	· · · · · · · · · · · · · · · · · · ·
SAG0220	92	
SAG0221	76	
SAG0222	108	
SAG0223	209	
SAG0224	332	
SAG0225	144	
SAG0226	418	
SAG0227	156	
SAG0228	111	
SAG0229	95	
SAG0229	96	
SAG0230	135	
SAG0231	186	
SAG0232	226	
SAG0234		
SAG0235		
SAG0236		
SAG0237	34	hypothetical protein



ORF	Size (a.a.)	Annotation	
SAG0238	41	hypothetical protein	
SAG0239	286	transcriptional regulator MutR family	
SAG0240	393	transporter, putative	
SAG0241	213	amino acid ABC transporter, permease protein	
SAG0242	308	amino acid ABC transporter, amino acid-binding protein	
SAG0243	211	amino acid ABC transporter, permease protein	
SAG0244	381	amino acid ABC transporter, ATP-binding protein	
SAG0245	152	protein of unknown function/lipoprotein, putative	
SAG0246	268	hypothetical protein	
SAG0247	116	hypothetical protein	
SAG0248	90	hypothetical protein	
SAG0249	116	hypothetical protein	
SAG0250	193	membrane protein, putative .	
SAG0251	72	transcriptional regulator, Cro/CI family	
SAG0252	186	acetyltransferase, GNAT family	
SAG0253	192	acetyltransferase, GNAT family	
SAG0254	226	acetyltransferase, GNAT family	
SAG0255	315	conserved hypothetical protein	
SAG0256	163	RNA polymerase sigma factor, ECF subfamily	
SAG0257	53	lipoprotein, putative	
SAG0258	202	transcriptional regulator, TetR family	
SAG0259	365	ABC transporter efflux protein, DrrB family, putative	
SAG0260	238	ABC transporter, ATP-binding protein	
SAG0261	129	IS1381, transposase OrfB	
SAG0262	127	IS1381, transposase OrfA	
SAG0263	171	hypothetical protein	
SAG0264	103	conserved hypothetical protein	
SAG0265	235		
SAG0266	382		
SAG0267	180	conserved hypothetical protein	
SAG0268	304		
SAG0269	213	acyl carrier protein phosphodiesterase, putative	
SAG0270	679		
SAG0271	85		
SAG0272	87		
SAG0273	502		
SAG0274	609		
SAG0275	232		
SAG0276	445		
SAG0277	476		
SAG0278	661		
SAG0279	101		
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SAG0285	298		

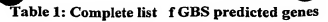
ORF	Size	Annotati n
Old	(a.a.)	
SAG0286	108	cell division protein FtsL, putative
SAG0287	752	penicillin-binding protein 2X
SAG0288	336	phospho-N-acetylmuramoyl-pentapeptide-transferase
SAG0289	447	ATP-dependent RNA helicase, DEAD/DEAH box family
SAG0290	270	ABC transporter, substrate-binding protein
SAG0291	267	amino acid ABC transporter, permease protein
SAG0292	247	amino acid ABC transporter, ATP-binding protein
SAG0293	74	conserved hypothetical protein
SAG0294	304	thioredoxin reductase
SAG0295	486	conserved hypothetical protein
SAG0296	273	NAD synthetase
SAG0297	444	aminopeptidase C
SAG0298	750	penicillin-binding protein 1A
SAG0299	199	recombination protein U
SAG0300	172	conserved hypothetical protein
SAG0301	40	hypothetical protein
SAG0302	110	conserved hypothetical protein
SAG0303	384	conserved hypothetical protein
SAG0304	487	conserved hypothetical protein
SAG0305	160	autoinducer-2 production protein LuxS
SAG0306	535	KH domain protein
SAG0307	33	hypothetical protein
SAG0308	298	ABC transporter, ATP-binding protein
SAG0309	246	ABC transporter, permease protein, putative
SAG0310	361	conserved hypothetical protein
SAG0311	NA	DNA-binding response regulator, authentic point mutation
SAG0312	234	conserved hypothetical protein
SAG0313	209	guanylate kinase
SAG0314	104	DNA-directed RNA polymerase, omega subunit, putative
SAG0315	796	
SAG0316	311	methionyl-tRNA formyltransferase
SAG0317	440	sun protein
SAG0318	245	serine/threonine phosphatase, putative
SAG0319	651	serine/threonine protein kinase
SAG0320	231	conserved hypothetical protein
SAG0321	339	· · · · · · · · · · · · · · · · · · ·
SAG0322	213	
SAG0323	466	<u></u>
		isomerase, cyclophilin type
SAG0324	124	<u></u>
SAG0325	258	
SAG0326	251	transcriptional regulator, DeoR family
SAG0327	327	
SAG0328	107	
SAG0329	106	_ <del> </del>
SAG0330	433	
SAG0331	818	
		· · · · · · · · · · · · · · · · · · ·



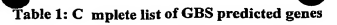
ORF	Size	Annotation
0.00000	(a.a.)	al1 Jahada sanas
SAG0333	362	glycerol dehydrogenase
SAG0334	308	cysteine synthase A
SAG0335	214	conserved hypothetical protein TIGR00257
SAG0336	429	helicase, putative
SAG0337	221	competence protein F, putative
SAG0338	184	ribosomal subunit interface protein
SAG0339	450	aspartate kinase family protein
SAG0340	216	hydrolase, haloacid dehalogenase-like family
SAG0341	49	hypothetical protein
SAG0342	263	enoyl-CoA hydratase/isomerase family protein
SAG0343	144	transcriptional regulator, MarR family
SAG0344	323	3-oxoacyl-(acyl-carrier-protein) synthase III
SAG0345	74	acyl carrier protein
SAG0346	319	enoyl-(acyl-carrier-protein) reductase II
SAG0347	308	malonyl CoA-acyl carrier protein transacylase
SAG0348	244	3-oxoacyl-[acyl-carrier protein] reductase
SAG0349	410	3-oxoacyl-(acyl-carrier-protein) synthase II
SAG0350	166	acetyl-CoA carboxylase, biotin carboxyl carrier protein
SAG0351	140	(3R)-hydroxymyristoyl-(acyl-carrier-protein) dehydratase
SAG0352	456	acetyl-CoA carboxylase, biotin carboxylase
SAG0353	291	acetyl-CoA carboxylase, carboxyl transferase, beta subunit
SAG0354	. 257	acetyl-CoA carboxylase, carboxyl transferase, alpha subunit
SAG0355	210	conserved hypothetical protein
SAG0356	425	seryl-tRNA synthetase
SAG0357	330	<u> </u>
SAG0358	120	<u></u>
SAG0359	303	PTS system, mannose-specific IID component
SAG0360	270	
SAG0361	336	<u> </u>
SAG0362	270	· · · · · · · · · · · · · · · · · · ·
SAG0363	194	<del></del>
SAG0364	203	
SAG0365	473	xanthine/uracil permease family protein
SAG0366	169	· · · · · · · · · · · · · · · · · · ·
SAG0367	186	
SAG0368	435	
SAG0369	98	· • · · · · · · · · · · · · · · · · · ·
SAG0309	139	
SAG0370	167	
SAG0371	85	
SAG0372 SAG0373	241	
SAG0373	344	
SAG0374 SAG0375		
	266	
SAG0376	211	
SAG0377	127	
SAG0378	379	
SAG0379	98	
SAG0380	100	ribosomal protein L7A family



	~-	
ORF	Size	Annotati n
G A G0201	(a.a.)	1
SAG0381	927	translation initiation factor IF-2
SAG0382	122	ribosome-binding factor A
SAG0383	334	protein of unknown function/lipoprotein, putative
SAG0384	138	transcriptional repressor CopY
SAG0385	744	copper-transporter ATPase CopA
SAG0386	68	copper-transporter protein CopZ
SAG0387	204	membrane protein, putative
SAG0388	270	hydrolase, haloacid dehalogenase-like family
SAG0389	880	DNA polymerase I
SAG0390	146	CoA-binding domain protein
SAG0391	159	transcriptional regulator, Fur family
SAG0392	521	cell wall surface anchor family protein
SAG0393	228	DNA-binding response regulator
SAG0394	345	sensor histidine kinase
SAG0395	246	membrane protein, putative
SAG0396	380	queuine tRNA-ribosyltransferase
SAG0397	102	conserved hypothetical protein
SAG0398	179	BioY family protein
SAG0399	258	AtsA/ElaC family protein
SAG0400	168	cytidine/deoxycytidylate deaminase family protein
SAG0401	44	hypothetical protein
SAG0402	449	glucose-6-phosphate isomerase
SAG0403	175	5-formyltetrahydrofolate cyclo-ligase family protein
SAG0404	225	rhomboid family protein
SAG0405	347	protein of unknown function/lipoprotein, putative
SAG0406	299	UTP-glucose-1-phosphate uridylyltransferase
SAG0407	338	glycerol-3-phosphate dehydrogenase (NAD(P)+)
SAG0408	109	ribonuclease P protein component
SAG0409	271	SpoIIIJ family protein
SAG0410	273	R3H domain protein
SAG0411	177	conserved hypothetical protein
SAG0412	258	recX protein '
SAG0413	451	RNA methyltransferase, TrmA family
SAG0414	153	conserved hypothetical protein
SAG0415	142	acetyltransferase, GNAT family
SAG0416	1233	protease, putative
SAG0417	302	glycosyl transferase, group 2 family protein
SAG0418	336	
SAG0419	137	nrdI protein
SAG0420	721	ribonucleoside-diphosphate reductase 2, alpha subunit
SAG0421	1055	
SAG0422	129	·
SAG0423	132	<u> </u>
SAG0424	94	
SAG0425	105	
SAG0426	131	conserved hypothetical protein
SAG0427	129	
SAG0428	345	



ORF	Size	Annotati n
21.50.400	(a.a.)	
SAG0429	284	oxidoreductase, aldo/keto reductase ramily
SAG0430	287	cation efflux system protein
SAG0431	174	transcriptional regulator, TetR family
SAG0432	397	transcriptional regulator, AraC family
SAG0433	1389	surface protein Rib
SAG0434	61	transposase, IS256 family, truncation
SAG0435	97	DNA-damage-inducible protein J, putative
SAG0436	62	hypothetical protein
SAG0437	123	lipoprotein, putative
SAG0438	145	bacteriophage L54a, integrase, truncation
SAG0439	NA	conserved hypothetical protein, degenerate
SAG0440	84	conserved hypothetical protein
SAG0441	103	conserved domain protein
SAG0442	189	acetyltransferase, GNAT family
SAG0443	194	acetyltransferase, GNAT family
SAG0444	188	conserved hypothetical protein
SAG0445	883	valyl-tRNA synthetase
SAG0446	319	oxidoreductase, Gfo/Idh/MocA family
SAG0447	287	magnesium transporter, CorA family
SAG0448	391	transposase, IS256 family
SAG0449	354	conserved hypothetical protein
SAG0450	330	aspartateammonia ligase
SAG0451	149	bacteriocin transport accessory protein, putative
SAG0452	179	type II DNA modification methyltransferase, putative
SAG0453	96	hypothetical protein
SAG0454	161	phosphopantetheine adenylyltransferase
SAG0455	357	conserved hypothetical protein
SAG0456	NA	conserved hypothetical protein, degenerate
SAG0457	192	conserved hypothetical protein
SAG0458	368	conserved hypothetical protein TIGR00048
SAG0459	171	VanZF domain protein
SAG0460	581	ABC transporter, ATP-binding/permease protein
SAG0461	579	ABC transporter, ATP-binding/permease protein
SAG0462	188	<u> </u>
SAG0463	179	·
SAG0464	330	·
SAG0465	164	
SAG0466	371	thiolase
SAG0467	409	·
SAG0468	210	· · · · · · · · · · · · · · · · · · ·
SAG0469	131	type IV prepilin peptidase-related protein
SAG0409	69	
SAG0470	322	
SAG0471		<u> </u>
	126	
SAG0473	613	
SAG0474	81	conserved hypothetical protein
SAG0475	451	UDP-N-acetylmuramoylalanineD-glutamate ligase
SAG0476	358	UDP-N-acetylglucosamineN-acetylmuramyl-(pentapeptide)

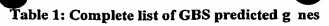


ORF	Size (a.a.)	Annotation
	(3.1.1)	pyrophosphoryl-undecaprenol N-acetylglucosamine transferase
SAG0477	378	cell division protein DivIB, putative
SAG0478	429	cell division protein FtsA
SAG0479	426	cell division protein FtsZ
SAG0480	224	ylmE protein, putative
SAG0481	201	ylmF protein
SAG0482	84	YGGT family protein
SAG0483	262	ylmH protein
SAG0484	256	cell division protein DivIVA, putative
SAG0485	930	isoleucyl-tRNA synthetase
SAG0486	100	conserved hypothetical protein
SAG0487	151	MutT/nudix family protein
SAG0488	753	ATP-dependent Clp protease, ATP-binding subunit
SAG0489	34	hypothetical protein
SAG0499	76	conserved hypothetical protein
SAG0490 SAG0491	230	amino acid ABC transporter, permease protein
SAG0491 SAG0492	244	amino acid ABC transporter, permease protein
SAG0492 SAG0493	564	
SAG0493 SAG0494	284	methylenetetrahydrofolate
5AG0494	204	dehydrogenase/methenyltetrahydrofolate cyclohydrolase
SAG0495	278	protein of unknown function
SAG0495		exodeoxyribonuclease VII, large subunit
	446 71	exodeoxyribonuclease VII, small subunit
SAG0497	290	geranyltranstransferase, putative
SAG0498	275	
SAG0499	157	
SAG0500		<del></del>
SAG0501	552	
SAG0502	278	DegV family protein lipase/acylhydrolase
SAG0503	279	
SAG0504	200	
SAG0505	91	DNA-binding protein HU
SAG0506	65	
SAG0507	310	
SAG0508	411	
SAG0509	403	
SAG0510	406	
SAG0511	270	
SAG0512	438	
SAG0513	128	
SAG0514		
SAG0515		
SAG0516		
SAG0517	374	
SAG0518	NA	
SAG0519		
SAG0520		
SAG0521	236	
SAG0522	232	metallo-beta-lactamase superfamily protein



ORF	Size (a.a.)	Annotation
SAG0523	254	oxidoreductase, short chain dehydrogenase/reductase family
SAG0524	835	DNA polymerase III, epsilon subunit/ATP-dependent helicase DinG
SAG0525	397	aspartate aminotransferase
SAG0526	448	asparaginyl-tRNA synthetase
SAG0527	185	conserved hypothetical protein
SAG0528	327	inosine-uridine preferring nucleoside hydrolase
SAG0529	38	hypothetical protein
SAG0530	137	OsmC/Ohr family protein
SAG0531	296	conserved hypothetical protein
SAG0532	324	conserved hypothetical protein
SAG0533	303	conserved hypothetical protein
SAG0534	465	dipeptidase
SAG0535	506	zinc ABC transporter, zinc-binding adhesion liprotein
SAG0536	86	ribosomal protein L31
SAG0537	311	DHH family protein
SAG0538	340	adenosine deaminase, putative
SAG0539	147	flavodoxin
SAG0540	91	chorismate mutase, putative
SAG0541	398	voltage-gated chloride channel family protein
SAG0542	127	IS1381, transposase OrfA
SAG0543	129	IS1381, transposase OrfB
SAG0544	115	ribosomal protein L19
SAG0545	359	prophage LambdaSa1, site-specific recombinase, phage integrase family
SAG0546	67	conserved domain protein
SAG0547	185	hypothetical protein
SAG0548	265	prophage LambdaSa1, repressor protein, putative
SAG0549	47	hypothetical protein
SAG0550	74	conserved hypothetical protein
SAG0551	52	conserved hypothetical protein
SAG0552	62	hypothetical protein
SAG0553	268	hypothetical protein
SAG0554	63	prophage LambdaSa1, transcriptional regulator, Cro/CI family
SAG0555	249	
SAG0556	47	hypothetical protein
SAG0557	76	<u> </u>
SAG0558	74	· · · · · · · · · · · · · · · · · · ·
SAG0559	286	<del></del>
SAG0560	77	
SAG0561	46	<u> </u>
SAG0562	84	
SAG0563	53	
SAG0564	160	<u>,                                     </u>
SAG0565	224	
SAG0566	138	<del></del>
SAG0567	439	
		protein

ORF	Size	Ann tation
	(a.a.)	TAMA LALIVIL
SAG0568	67	conserved hypothetical protein
SAG0569	158	conserved hypothetical protein
SAG0570	115	hypothetical protein
SAG0571	43	hypothetical protein
SAG0572	138	conserved hypothetical protein
SAG0573	54	hypothetical protein
SAG0574	89	conserved hypothetical protein
SAG0575	110	hypothetical protein
SAG0576	43	hypothetical protein
SAG0577	177	conserved hypothetical protein
SAG0577	88	conserved hypothetical protein
SAG0579	142	conserved hypothetical protein
SAG0580	111	conserved hypothetical protein, truncation
SAG0580	118	conserved hypothetical protein
SAG0582	422	conserved hypothetical protein
SAG0583	406	conserved hypothetical protein
SAG0584	62	conserved hypothetical protein, truncation
SAG0585	471	conserved hypothetical protein
SAG0586	154	conserved hypothetical protein
SAG0587	300	prophage LambdaSa1, structural protein, putative
SAG0588	71	conserved hypothetical protein
SAG0589	143	conserved hypothetical protein
SAG0590	112	conserved hypothetical protein
SAG0591	78	conserved hypothetical protein
SAG0592	111	conserved hypothetical protein
SAG0593	185	prophage LambdaSa1, structural protein
SAG0594	81	conserved hypothetical protein
SAG0595	123	conserved hypothetical protein
SAG0596	670	prophage LambdaSal, pblA protein, internal deletion
SAG0597	506	prophage LambdaSa1, minor structural protein, putative
SAG0598	1374	prophage LambdaSa1, N-acetylmuramoyl-L-alanine amidase,
		family 4
SAG0599	668	prophage LambdaSa1, minor structural protein, putative
SAG0600	109	
SAG0601	70	hypothetical protein
SAG0602	100	<u> </u>
SAG0603	111	conserved hypothetical protein
SAG0604	239	prophage LambdaSa1, lysin, putative
SAG0605	323	conserved hypothetical protein
SAG0606	66	conserved hypothetical protein
SAG0607	56	
SAG0608	59	
SAG0609	NA	prophage LambdaSa1, integrase, degenerate
SAG0610	134	
SAG0611	NA	transposase, degenerate
SAG0612	53	conserved hypothetical protein
SAG0613	425	transmembrane protein Vexp1
SAG0614	218	

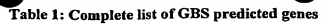


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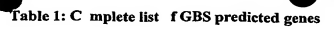
ORF	Siz	Ann totion
		Ann tation
SAG0662	(a.a.) 101	cylX protein
SAG0663	282	cylD protein
SAG0664	240	cylG protein
SAG0665	101	acyl carrier protein AcpC
SAG0666	158	cylZ protein
SAG0667	309	cylA protein
SAG0668	292	cylB protein
SAG0669	667	cylE protein
SAG0670	317	cylF protein
SAG0671	731	cyll protein
SAG0672	403	cylJ protein
SAG0673	191	cylK protein
SAG0674	113	hypothetical protein
SAG0675	171	putative secreted protein
SAG0676	885	proteinase, putative
SAG0677	1062	hypothetical protein
SAG0678	NA	endopeptidase O, degenerate
SAG0679	343	protein of unknown function
SAG0680	339	protein of unknown function
SAG0681	353	conserved domain protein
SAG0682	409	permease, putative
SAG0683	NA	transmembrane protein Vexp3, putative, degenerate
SAG0684	223	ABC transporter, ATP-binding protein
SAG0685	472	conserved hypothetical protein
SAG0686	261	DNA-entry nuclease, putative
SAG0687	212	DedA family protein, putative
SAG0688	218	ABC transporter, ATP-binding protein
SAG0689	257	membrane protein, putative
SAG0690	272	conserved hypothetical protein
SAG0691	294	transcriptional regulator, LysR family
SAG0692	193	regulatory protein, putative
SAG0693	377	IS1548, transposase
SAG0694	173	regulatory protein, putative, truncation
SAG0695	330	D-lactate dehydrogenase
SAG0696	516	sodium:galactoside symporter family protein, putative
SAG0697	341	2-keto-3-deoxygluconate kinase
SAG0698	599	beta-glucuronidase
SAG0699	223	transcriptional regulator, GntR family
SAG0700	205	2-dehydro-3-deoxyphosphogluconate aldolase/4-hydroxy-2-
		oxoglutarate aldolase
SAG0701	466	glucuronate isomerase
SAG0702	348	mannonate dehydratase
SAG0703	279	
SAG0704	270	<u> </u>
SAG0705	596	glycosyl hydrolase, family 3
SAG0706	361	proline dipeptidase
SAG0707	334	transcriptional regulator, RegM family
SAG0708	488	alpha amylase family protein

ORF	Size	Ann tati n
	(a.a.)	
SAG0709	332	glycosyl transferase, group 1 family protein
SAG0710	444	glycosyl transferase, group 1 family protein
SAG0711	647	threonyl-tRNA synthetase
SAG0712	234	DNA-binding response regulator
SAG0713	339	conserved hypothetical protein
SAG0714	188	conserved hypothetical protein
SAG0715	216	amino acid ABC transporter, permease protein
SAG0716	231	amino acid ABC transporter, permease protein
SAG0717	266	amino acid ABC transporter, amino acid-binding protein
SAG0718	251	amino acid ABC transporter, ATP-binding protein
SAG0719	236	DNA-binding response regulator
SAG0720	449	sensory box histidine kinase
SAG0721	269	metallo-beta-lactamase superfamily protein
SAG0722	122	conserved hypothetical protein
SAG0723	236	ribonuclease III
SAG0724	1179	chromosome segregation SMC protein
SAG0725	265	hydrolase, haloacid dehalogenase-like family
SAG0726	274	hydrolase, haloacid dehalogenase-like family
SAG0727	536	signal recognition particle-docking protein FtsY
SAG0728	270	ABC transporter, substrate-binding protein
SAG0729	300	ABC transporter, permease protein, putative
SAG0730	42	ABC transporter, ATP-binding protein
SAG0731	347	bacterial luciferase family protein
SAG0732	720	transcriptional accessory protein Tex, putative
SAG0733	142	conserved hypothetical protein
SAG0734	87	phage shock protein C, putative
SAG0735	44	hypothetical protein
SAG0736	311	HPr(Ser) kinase/phosphatase
SAG0737	257	prolipoprotein diacylglyceryl transferase
SAG0738	132	conserved hypothetical protein
SAG0739	143	conserved hypothetical protein
SAG0740	91	conserved hypothetical protein
SAG0741	303	peptidase, U32 family, putative
SAG0742	428	peptidase, U32 family
SAG0743	70	conserved hypothetical protein
SAG0744	265	
SAG0745	446	
SAG0746	369	<del></del>
SAG0747	208	· · · · · · · · · · · · · · · · · · ·
SAG0748	397	·
SAG0749	156	
SAG0750	496	<del></del>
SAG0751	300	hydrolase, haloacid dehalogenase-like family
SAG0752	213	phosphoglycerate mutase family protein
SAG0753	157	ebsC family protein, putative
SAG0754	205	+ <del></del>
SAG0755	282	peptidase, U32 family
SAG0756	174	conserved hypothetical protein

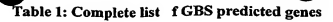


ORF	Size (a.a.)	Annotation
SAG0757	129	protein of unknown function/lipoprotein, putative
SAG0758	599	oligoendopeptidase F, putative
SAG0759	931	phosphoenolpyruvate carboxylase
SAG0760	377	IS1548, transposase
SAG0761	422	cell division protein, FtsW/RodA/SpoVE family
SAG0762	398	translation elongation factor Tu
SAG0763	252	triosephosphate isomerase
SAG0764	230	phosphoglycerate mutase family protein
SAG0765	681	penicillin-binding protein 2b
SAG0766	198	recombination protein RecR
SAG0767	348	D-alanine-D-alanine ligase
SAG0768	455	UDP-N-acetylmuramoylalanyl-D-glutamyl-2,6-diaminopimelate-
02100700	"55	D-alanyl-D-alanyl ligase
SAG0769	406	oxalate:formate antiporter
SAG0770	228	membrane protein, putative
SAG0770	512	cell wall surface anchor family protein
SAG0771	514	peptide chain release factor 3
SAG0773	126	conserved hypothetical protein
SAG0773	244	ABC transporter, ATP-binding protein
SAG0774 SAG0775	+	ABC transporter, permease protein
<del></del>	220	
SAG0776	276	YaeC family protein, putative
SAG0777	528	ATP-dependent RNA helicase, DEAD/DEAH box family
SAG0778	88	conserved hypothetical protein
SAG0779	254	conserved hypothetical protein
SAG0780	246	acyltransferase family protein
SAG0781	217	competence protein CelA
SAG0782	745	DNA internalization-related competence protein ComEC/Rec2
SAG0783	269	hydrolase, haloacid dehalogenase-like family
SAG0784	314	sugar-binding transcriptional regulator, LacI family
SAG0785	330	
SAG0786	242	
SAG0787	345	
SAG0788	202	
SAG0789	283	
SAG0790	622	
SAG0791	475	
SAG0792	364	
SAG0793	380	
SAG0794	418	
SAG0795	354	
SAG0796	147	
SAG0797	342	
SAG0798	226	
SAG0799	233	glucosamine-6-phosphate isomerase
-SAG0800	318	
SAG0801	239	ribosomal small subunit pseudouridine synthase A
SAG0802	38	
SAG0803	383	major facilitator family protein

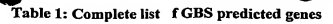
ODE	C:	
ORF	Size	Annotation
SAG0804	(a.a.)	Compotence matrix Coi A
		competence protein CoiA
SAG0805	601	oligoendopeptidase B
SAG0806	208	hydrolase, haloacid dehalogenase-like family
SAG0807	235	O-methyltransferase family protein
SAG0808	309	protease maturation protein, putative
SAG0809	161	conserved hypothetical protein
SAG0810	872	alanyl-tRNA synthetase
SAG0811	238	membrane protein, putative
SAG0812	272	glycosyl transferase, family 8
SAG0813	81	hypothetical protein .
SAG0814	95	conserved hypothetical protein
SAG0815	71	transcriptional regulator, Cro/CI family
SAG0816	253	membrane protein, putative
SAG0817	187	membrane protein, putative
SAG0818	319	ribonucleoside-diphosphate reductase 2, beta subunit
SAG0819	719	ribonucleoside-diphosphate reductase 2, alpha subunit
SAG0820	74	ribonucleoside-diphosphate reductase 2, NrdH-redoxin
SAG0821	87	phosphocarrier protein HPr
SAG0822	577	phosphoenolpyruvate-protein phosphotransferase
SAG0823	475	glyceraldehyde-3-phosphate dehydrogenase, NADP-dependent
SAG0824	417	polysaccharide deacetylase family protein
SAG0825	360	ATP-dependent RNA helicase, DEAD/DEAH box family
SAG0826	209	uridine kinase
SAG0827	165	conserved hypothetical protein
SAG0828	554	DNA polymerase III, gamma and tau subunits
SAG0829	64	conserved hypothetical protein
SAG0830	311	biotinacetyl-CoA-carboxylase ligase
SAG0831	398	S-adenosylmethionine synthetase
SAG0832	753	protein of unknown function
SAG0833	181	hypothetical protein
SAG0834	42	hypothetical protein
SAG0835	188	conserved hypothetical protein
SAG0836	184	conserved hypothetical protein
SAG0837	428	
SAG0838	233	· · · · · · · · · · · · · · · · · · ·
SAG0839	226	<u> </u>
SAG0840	265	+ · · · · <u> </u>
SAG0841	256	
SAG0842	223	thiamine-phosphate pyrophosphorylase
SAG0843	419	
SAG0844	184	<u> </u>
SAG0845	427	·
SAG0846	286	· · · · · · · · · · · · · · · · · · ·
SAG0847	306	
SAG0847	151	
SAG0849		GtrA family protein
	169	
SAG0850	652	<u> </u>
SAG0851	339	bmrU protein, putative



ODE	C:	
ORF	Size	Annotation
04.00060	(a.a.)	nullulanace nutotive
SAG0852	766	partitionase, putative
SAG0853	622	1,4-alpha-glucan branching enzyme
SAG0854	379	glucose-1-phosphate adenylyltransferase
SAG0855	NA	glycogen biosynthesis protein GlgD, authentic frameshift
SAG0856	476	glycogen synthase
SAG0857	66	ATP synthase F0, C subunit
SAG0858	238	ATP synthase F0, A subunit
SAG0859	165	ATP synthase F0, B subunit
SAG0860	178	ATP synthase F1, delta subunit
SAG0861	501	ATP synthase F1, alpha subunit
SAG0862	293	ATP synthase F1, gamma subunit
SAG0863	468	ATP synthase F1, beta subunit
SAG0864	137	ATP synthase F1, epsilon subunit
SAG0865	76	conserved hypothetical protein
SAG0866	423	UDP-N-acetylglucosamine 1-carboxyvinyltransferase
SAG0867	63	conserved hypothetical protein
SAG0868	285	DNA-entry nuclease
SAG0869	346	phenylalanyl-tRNA synthetase, alpha subunit
SAG0870	173	acetyltransferase, GNAT family
SAG0871	801	phenylalanyl-tRNA synthetase, beta subunit
SAG0872	300	conserved hypothetical protein
SAG0873	1077	exonuclease RexB
SAG0874	1207	exonuclease RexA
SAG0875	305	magnesium transporter, CorA family, putative
SAG0876	458	tRNA modification GTPase TrmE
SAG0877	636	ABC transporter, ATP-binding protein
SAG0878	322	acetoin dehydrogenase, thymine PPi dependent, E1 component,
		alpha subunit
SAG0879	332	acetoin dehydrogenase, thymine PPi dependent, E1 component,
		beta subunit
SAG0880	462	acetoin dehydrogenase, thymine PPi dependent, E2 component,
		dihydrolipoamide acetyltransferase
SAG0881	585	acetoin dehydrogenase, thymine PPi dependent, E3 component,
		dihydrolipoamide dehydrogenase
SAG0882	329	lipoate-protein ligase A
SAG0883	261	cobyric acid synthase, putative
SAG0884	447	mur ligase family protein
SAG0885	283	conserved hypothetical protein TIGR00159
SAG0886	319	protein of unknown function
SAG0887	450	phosphoglucomutase/phosphomannomutase family protein
SAG0888	123	conserved hypothetical protein
SAG0889	126	conserved hypothetical protein
SAG0890	376	oxygen-independent coproporphyrinogen III oxidase, putative
SAG0891	245	conserved hypothetical protein
SAG0892	256	hydrolase, haloacid dehalogenase-like family
SAG0893	218	conserved hypothetical protein
SAG0894	1370	protein of unknown function
SAG0895	289	
DAG0033	1 209	lipoyl-binding domain protein

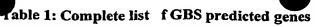


SAG0896 108 oxidoreductase, putative SAG0897 221 conserved hypothetical protein SAG0898 83 hypothetical protein SAG0899 57 hypothetical protein SAG0900 56 hypothetical protein SAG0901 127 hypothetical protein SAG0902 45 hypothetical protein SAG0903 44 hypothetical protein SAG0904 56 hypothetical protein SAG0905 138 nucleoside diphosphate kinase SAG0906 610 GTP-binding protein LepA SAG0907 877 protein of unknown function/lipoprotein, putative SAG0908 203 HD domain protein SAG0909 154 acetyltransferase, GNAT family SAG0910 144 PilB-related protein SAG0911 930 cation-transporting ATPase, E1-E2 family SAG0913 212 chloramphenicol acetyltransferase SAG0914 203 conserved hypothetical protein SAG0915 405 Tn916, transposase SAG0916 67 Tn916, excisionase SAG0917 83 Tn916, hypothetical protein	
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SAG0914 203 conserved hypothetical protein SAG0915 405 Tn916, transposase SAG0916 67 Tn916, excisionase SAG0917 83 Tn916, hypothetical protein	
SAG0915         405         Tn916, transposase           SAG0916         67         Tn916, excisionase           SAG0917         83         Tn916, hypothetical protein	
SAG0916 67 Tn916, excisionase SAG0917 83 Tn916, hypothetical protein	
SAG0917 83 Tn916, hypothetical protein	
SAG0918 76 Tn916, hypothetical protein	
SAG0919   157   Tn916, hypothetical protein	
SAG0920 23 Tn916, hypothetical protein	
SAG0921 117 Tn916, transcriptional regulator, putative	
SAG0922 61 Tn916, hypothetical protein	
SAG0923 639 Tn916, tetracycline resistance protein	
SAG0924 28 Tn916, tetM leader peptide	
SAG0925 310 Tn916, hypothetical protein	
SAG0926 333 Tn916, NLP/P60 family protein	
SAG0927 725 membrane protein, putative	
SAG0928 NA Tn916, hypothetical protein, authentic frameshift	
SAG0929 168 Tn916, hypothetical protein	
SAG0930 165 Tn916, hypothetical protein	
SAG0931 73 Tn916, hypothetical protein	
SAG0932 401 Tn916, transcriptional regulator, putative	
SAG0933 461 Tn916, FtsK/SpoIIIE family protein	
SAG0934 128 Tn916, hypothetical protein	
SAG0935 104 Tn916, hypothetical protein	
SAG0936 39 Tn916, hypothetical protein	
SAG0937 NA ABC transporter, ATP-binding protein, authentic framesh	ift
SAG0938 122 transcriptional regulator, GntR family	
SAG0939 1034 DNA polymerase III, alpha subunit	
SAG0940 340 6-phosphofructokinase	
SAG0941 500 pyruvate kinase	
SAG0942 185 signal peptidase I, putative	
SAG0943 47 hypothetical protein	

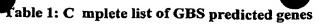


ORF	Size	Annotation
	(a.a.)	
SAG0944	604	glucosaminefructose-6-phosphate aminotransferase, isomerizing
SAG0945	377	IS1548, transposase
SAG0946	109	phnA protein
SAG0947	213	amino acid ABC transporter, permease protein
SAG0948.	209	amino acid ABC transporter, ATP-binding protein
SAG0949	276	amino acid ABC transporter, amino acid-binding protein
SAG0950	82	ribosomal protein S20
SAG0951	306	pantothenate kinase
SAG0952	196	conserved hypothetical protein
SAG0953	129	cytidine deaminase
SAG0954	. 349	protein of unknown function/lipoprotein, putative
SAG0955	511	sugar ABC transporter, ATP-binding protein
SAG0956	353	sugar ABC transporter, permease protein, putative
SAG0957	318	sugar ABC transporter, permease protein, putative
SAG0958	456	NADH oxidase
SAG0959	329	L-lactate dehydrogenase
SAG0960	819	DNA gyrase, A subunit
SAG0961	247	sortase SrtA
SAG0962	137	glyoxylase family protein
SAG0963	320	conserved hypothetical protein
SAG0964	375	Na+/H+ exchanger family protein
SAG0965	127	IS1381, transposase OrfA
SAG0966	129	IS1381, transposase OrfB
SAG0967	520	GMP synthase
SAG0968	232	transcriptional regulator, GntR family
SAG0969	444	gid protein
SAG0970	247	acetyltransferase, GNAT family
SAG0971	282	protein of unknown function/lipoprotein, putative
SAG0972	NA	conserved hypothetical protein, authentic frameshift
SAG0973	320	nisin-resistance protein, putative
SAG0974	250	ABC transporter, ATP-binding protein
SAG0975	651	ABC transporter, permease protein, putative
SAG0976	222	DNA-binding response regulator
SAG0977	312	sensor histidine kinase
SAG0978	356	
SAG0979	553	ABC transporter, substrate-binding protein
SAG0980	257	conserved hypothetical protein
SAG0981	228	satD protein
SAG0982	521	signal recognition particle protein Ffh
SAG0983	110	conserved hypothetical protein
SAG0984	437	sensor histidine kinase CiaH
SAG09°5	226	DNA-binding response regulator CiaR
SAG0986	849	aminopeptidase N
SAG0987	217	phosphate transport system regulatory protein PhoU
SAG0988	252	phosphate ABC transporter, ATP-binding protein PstB, putative
SAG0989	267	phosphate ABC transporter, ATP-binding protein PstB, putative
SAG0990	295	phosphate ABC transporter, permease protein PstA, putative
SAG0991	305	phosphate ABC transporter, permease protein

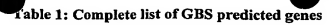
ODE	G:	
ORF	Siz (a.a.)	Annotati n
SAG0992	286	phosphate ABC transporter, phosphate-binding protein
SAG0993	436	NOL1/NOP2/sun family protein
SAG0994	254	inositol monophosphatase family protein
SAG0995	93	conserved hypothetical protein
SAG0996	137	conserved hypothetical protein
SAG0997	310	macrolide-efflux protein mreA/riboflavin biosynthesis protein RibF
SAG0998	294	tRNA pseudouridine synthase B
SAG0999	143	acetyltransferase, GNAT family
SAG1000	423	conserved hypothetical protein
SAG1001	196	conserved hypothetical protein
SAG1002	292	protease, putative
SAG1003	876	permease, putative
SAG1004	233	ABC transporter, ATP-binding protein
SAG1005	706	
SAG1006	280	DprA/SMF protein, putative DNA processing factor
SAG1007	342	iron-compound ABC transporter, iron-compound-binding protein
SAG1008	253	iron compound ABC transporter, ATP-binding protein
SAG1009	324	iron compound ABC transporter, permease protein
SAG1010	320	iron compound ABC transporter, permease protein
SAG1011	182	acetyltransferase, CysE/LacA/LpxA/NodL family
SAG1012	253	ribonuclease HII
SAG1013	283	GTP-binding protein
SAG1014	190	conserved hypothetical protein
SAG1015	494	carbon starvation protein CstA, putative
SAG1016	244	response regulator
SAG1017	579	sensor histidine kinase, putative
SAG1018	40	lipoprotein, putative
SAG1019	39	hypothetical protein
SAG1020	227	lipoprotein, putative
SAG1021	107	hypothetical protein
SAG1022	177	hypothetical protein
SAG1023	48	hypothetical protein
SAG1024	183	lipoprotein, putative
SAG1025	149	hypothetical protein
SAG1026	NA	immunogenic secreted protein, degenerate
SAG1027	84	conserved hypothetical protein
SAG1028	196	
SAG1029	101	hypothetical protein
SAG1029	304	protein of unknown function
SAG1031	120	conserved domain protein
SAG1031	85	conserved hypothetical protein
SAG1032	1309	
SAG1033		
SAG1034	55	
SAG1035 SAG1036	424	conserved hypothetical protein
	80	conserved hypothetical protein
SAG1037	157	hypothetical protein
SAG1038	1003	phage infection protein, putative



ORF	Size	Annotation
6.6466	(a.a.)	
SAG1039	96	conserved hypothetical protein
SAG1040	260	conserved domain protein
SAG1041	107	hypothetical protein
SAG1042	1060	carbamoyl-phosphate synthase, large subunit
SAG1043	358	carbamoyl-phosphate synthase, small subunit
SAG1044	307	aspartate carbamoyltransferase
SAG1045	430	dihydroorotase, multifunctional complex type
SAG1046	209	orotate phosphoribosyltransferase
SAG1047	233	orotidine 5'-phosphate decarboxylase
SAG1048 -	410	membrane protein, putative
SAG1049	513	ABC transporter, ATP-binding protein
SAG1050	112	ribonucleotide reductase, truncation
SAG1051	358	aspartate-semialdehyde dehydrogenase
SAG1052	47	cell wall surface anchor family protein, putative
SAG1053	30	hypothetical protein
SAG1054	531	cardiolipin synthetase
SAG1055	556	formatetetrahydrofolate ligase
SAG1056	339	lipoate-protein ligase A
SAG1057	292	conserved hypothetical protein
SAG1058	272	conserved hypothetical protein
SAG1059	110	glycine cleavage system H protein, putative
SAG1060	328	bacterial luciferase family protein
SAG1061	399	oxidoreductase, FMN-binding .
SAG1062	282	lipoate-protein ligase A family protein
SAG1063	228	flavoprotein-related protein
SAG1064	180	flavoprotein family protein
SAG1065	190	membrane protein, putative
SAG1066	572	phosphoglucomutase
SAG1067	178	IS861, transposase OrfA
SAG1068	277	IS861, transposase OrfB
SAG1069	65	hypothetical protein
SAG1070	577	ABC transporter, ATP-binding/permease protein
SAG1071	573	ABC transporter, ATP-binding/permease protein
SAG1072	200	conserved hypothetical protein
SAG1073	325	conserved hypothetical protein
SAG1074	418	serine hydroxymethyltransferase
SAG1075	183	Sua5/YciO/YrdC/YwlC family protein
SAG1076	276	modification methylase, HemK family
SAG1077	359	peptide chain release factor 1
SAG1078	189	thymidine kinases
SAG1079	60	4-oxalocrotonate tautomerase
SAG1080	47	hypothetical protein
SAG1081	312	ApbE family protein
SAG1082	200	conserved hypothetical protein
SAG1083	411	conserved hypothetical protein
SAG1084	262	formate/nitrite transporter family protein
SAG1085	424	xanthine permease
SAG1086	193	xanthine phosphoribosyltransferase

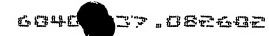


Size	Annotation
(a.a.)	
	guanosine monophosphate reductase
	drug resistance transporter, EmrB/QacA family, putative
230	conserved hypothetical protein
666	potassium uptake protein, putative
216	oxidoreductase, short chain dehydrogenase/reductase family
330	phosphate acetyltransferase
294	ribosomal large subunit pseudouridine synthase, RluD subfamily
278	conserved hypothetical protein
223	GTP pyrophosphokinase family protein
190	conserved hypothetical protein
324	ribose-phosphate pyrophosphokinase
371	cysteine desulphurase
	conserved hypothetical protein
	conserved hypothetical protein
	DNA repair protein RadC
	membrane protein, putative
	6-phospho-beta-glucosidase
	platelet activating factor, putative
	hydrolase, haloacid dehalogenase-like family
	transcriptional regulator, AraC family, putative
	voltage-gated chloride channel family protein
·	spermidine/putrescine ABC transporter, spermidine/putrescine-
337	binding protein
258	spermidine/putrescine ABC transporter, permease protein
	spermidine/putrescine ABC transporter, permease protein
	spermidine/putrescine ABC transporter, ATP-binding protein
	UDP-N-acetylenolpyruvoylglucosamine reductase
162	2-amino-4-hydroxy-6-hydroxymethyldihydropteridine
	pyrophosphokinase
120	dihydroneopterin aldolase
267	dihydropteroate synthase
187	GTP cyclohydrolase I
420	folylpolyglutamate synthase
295	
288	<del> </del>
	homoserine dehydrogenase
	<u> </u>
	aldehyde dehydrogenase family protein
	membrane protein, putative
	protein of unknown function
	conserved domain protein
	<del>                                     </del>
36	transcriptional regulator, Cro/CI family hypothetical protein
. 20	i mydomencai diolem
49 164	hypothetical protein thiol peroxidase
	(a.a.) 327 446 230 666 216 330 294 278 223 190 324 371 115 210 226 377 478 204 273 309 510 357  258 264 384 300 162 120 267 187 420 295 288 427 295 515 34 458 335 228 446 65



ORF	Size	Annotation
	(a.a.)	Anaotation
SAG1133	254	conserved hypothetical protein
SAG1134	213	transcriptional regulator, GntR family/potassioum uptake protein,
3.13.1		TrkA family
SAG1135	183	gls24 protein, putative
SAG1136	65	conserved hypothetical protein
SAG1137	180	gls24 protein, putative
SAG1138	64	conserved hypothetical protein
SAG1139	193	conserved hypothetical protein
SAG1140	82	conserved hypothetical protein
SAG1141	112	conserved hypothetical protein
SAG1142	759	ATP-dependent DNA helicase PcrA
SAG1142	128	conserved hypothetical protein
SAG1144	441	uracil permease
SAG1145	448	sodium:alanine symporter family protein
SAG1146	411	cation efflux family protein
SAG1147	130	
SAG1148	231	conserved hypothetical protein
SAG1148	207	membrane protein, putative
	<del>+</del>	lipoprotein, putative
SAG1150	400	ribosomal protein S1
SAG1151	76 340	conserved hypothetical protein
SAG1152		branched-chain amino acid aminotransferase
SAG1153	819	DNA topoisomerase IV, A subunit
SAG1154	653	DNA topoisomerase IV, B subunit
SAG1155	212	membrane protein, putative
SAG1156	217	uracil-DNA glycosylase
SAG1157	161	conserved hypothetical protein
SAG1158	413	CMP-N-acetylneuraminic acid synthetase NeuA
SAG1159	209	neuD protein
SAG1160	384	UDP-N-acetylglucosamine-2-epimerase NeuC
SAG1161	341	N-acetyl neuramic acid synthetase NeuB
SAG1162	466	polysaccharide biosynthesis protein CpsL
SAG1163	318	polysaccharide biosynthesis protein CpsK(V)
SAG1164	321	glycosyl transferase CpsJ(V)
SAG1165	327	glycosyl transferase CpsO(V)
SAG1166	295	
SAG1167	241	polysaccharide biosynthesis protein CpsM(V)
SAG1168	364	<u> </u>
SAG1169	163	
SAG1170	149	
SAG1171	462	
SAG1172	229	
SAG1173	230	
SAG1174	243	capsular polysaccharide biosynthesis protein CpsB
SAG1175	485	capsular polysaccharide biosynthesis protein CpsA
SAG1176	290	
SAG1177	255	conserved hypothetical protein
SAG1178	236	purine nucleoside phosphorylase
SAG1179	418	voltage-gated chloride channel family protein, putative

ORF	Size	Annotation
OKF	(a.a.)	Annotation
SAG1180	269	purine nucleoside phosphorylase
SAG1180	135	arsenate reductase
SAG1181	403	phosphopentomutase
SAG1182	223	ribose 5-phosphate isomerase
SAG1183	236	conserved hypothetical protein
SAG1185	262	tributyrin esterase
SAG1185	553	
SAG1180	253	metallo-beta-lactamase superfamily protein
SAG1188	287	ABC transporter, ATP-binding protein
SAG1189	334	ABC transporter, permease protein
	551	conserved hypothetical protein
SAG1190	239	adherence and virulence protein A
SAG1191		alpha-acetolactate decarboxylase
SAG1192	560	acetolactate synthase, catabolic
SAG1193	408	TPR domain protein
SAG1194	396	membrane protein, putative
SAG1195	153	MutT/nudix family protein
SAG1196 SAG1197	160 1072	mutator MutT protein
		hyaluronidase
SAG1198 SAG1199	348	dTDP-glucose 4,6-dehydratase
	197	dTDP-4-dehydrorhamnose 3,5-epimerase
SAG1200	289	glucose-1-phosphate thymidylyltransferase
SAG1201 SAG1202	367	iminodiacetate oxidase, putative
SAG1202 SAG1203	262	conserved hypothetical protein TIGR00486
	227	conserved hypothetical protein
SAG1204 SAG1205	226 172	DNA replication protein DnaD, putative
SAG1205	854	adenine phosphoribosyltransferase
SAG1200	32	conserved domain protein hypothetical protein
SAG1207 SAG1208	732	
SAG1208	253	single-stranded-DNA-specific exonuclease RecJ
SAG1210		oxidoreductase, short chain dehydrogenase/reductase family
SAG1210 SAG1211	309 215	metallo-beta-lactamase superfamily protein
SAG1211 SAG1212		conserved hypothetical protein
SAG1212 SAG1213	412	GTP-binding protein HflX
	296	tRNA delta(2)-isopentenylpyrophosphate transferase
SAG1214 SAG1215	58	hypothetical protein
SAG1215 SAG1216	305	exfoliative toxin A, putative
	1252	pullulanase, putative
SAG1217 SAG1218	NA 104	conserved hypothetical protein, authentic frameshift
	194	conserved hypothetical protein
SAG1219	468	peptidase, M20/M25/M40 family
SAG1220	200	nitroreductase family protein
SAG1221	NA	glycerophosphoryl diester phosphodiesterase, putative, authentic
GAG1222	500	point mutation
SAG1222	593	excinuclease ABC, C subunit
SAG1223	255	conserved hypothetical protein
SAG1224	446	MATE efflux family protein
SAG1225	136	conserved hypothetical protein
SAG1226	165	conserved hypothetical protein



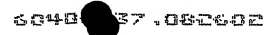
ORF	Size	Annotation
	(a.a.)	
SAG1227	198	protein of unknown function
SAG1228		ISSdy1, transposase OrfA
SAG1229_	259	
SAG1230	96	conserved hypothetical protein
SAG1231	NA	transposase OrfB, IS3 family, degenerate
SAG1232	77	transposase OrfB, IS3 family, truncation
SAG1233	822	streptococcal histidine triad family protein
SAG1234	306	laminin-binding surface protein
SAG1235	425	GBSi1, group II intron, maturase
SAG1236	NA	C5a peptidase, authentic frameshift
SAG1237	444	hypothetical protein
SAG1238	202	hypothetical protein
SAG1239	76	conserved hypothetical protein
SAG1240	125	conserved hypothetical protein, truncation
SAG1241	91	transposase OrfA, IS3 family
SAG1242	67	transposase OrfB, IS3 family, truncation
SAG1243	96	ISSdy1, transposase OrfA
SAG1244	259	ISSdy1, transposase OrfB
SAG1245	38	hypothetical protein
SAG1246	389	hypothetical protein .
SAG1247	399	site-specific recombinase, phage integrase family
SAG1248	75	conserved hypothetical protein
SAG1249	74	transcriptional regulator, Cro/CI family
SAG1250	621	Tn5252, relaxase
SAG1251	121	Tn5252, Orf 9 protein
SAG1252	120	Tn5252, Orf 10 protein
SAG1253	435	transposase, ISL3 family
SAG1254	546	mercuric reductase
SAG1255	130	mercuric resistance operon regulatory protein MerR
SAG1256	142	IS861, transposase OrfB, truncation
SAG1257	709	cation-transporting ATPase, E1-E2 family
SAG1258	122	cadmium efflux system accessory protein
SAG1259	99	conserved hypothetical protein
SAG1260	262	hypothetical protein
SAG1261	198	conserved hypothetical protein
SAG1262	695	cation-transporting ATPase, E1-E2 family
SAG1263	NA	conserved domain protein, authentic frameshift
SAG1264	148	transcriptional repressor CopY, putative
SAG1265	206	cadmium resistance transporter, putative
SAG1266	152	hypothetical protein
SAG1267	108	
SAG1268	230	repressor protein, putative
SAG1269	44	hypothetical protein
SAG1270	471	ImpB/MucB/SamB family protein
SAG1271	116	conserved hypothetical protein
SAG1272	102	conserved hypothetical protein
SAG1273	118	conserved hypothetical protein
SAG1274	129	conserved hypothetical protein

ORF	Size	Ann tation
	(a.a.)	Aun tauou
SAG1275	75	hypothetical protein
SAG1276	358	conserved hypothetical protein
SAG1277	163	hypothetical protein
SAG1278	96	hypothetical protein
SAG1279	99	conserved domain protein
SAG1280	2274	SNF2 family protein
SAG1281	183	hypothetical protein
SAG1282	63	calcium-binding protein, putative
SAG1283	1631	agglutinin receptor
SAG1284	196	abortive infection protein AbiGI
SAG1285	281	abortive infection protein AbiGII
SAG1286	933	Tn5252, Orf28
SAG1287	776	Tn5252, Orf26
SAG1288	NA	Tn5252, Orf25, degenerate
SAG1289	284	Tn5252, Orf23
SAG1290	80	hypothetical protein
SAG1291	605	Tn5252, Orf 21 protein, internal deletion
SAG1292	162	hypothetical protein
SAG1293	194	protease, putative
SAG1294	77	conserved hypothetical protein
SAG1295	127	conserved hypothetical protein
SAG1296	142	conserved hypothetical protein
SAG1297	451	C-5 cytosine-specific DNA methylase
SAG1298	31	hypothetical protein
SAG1299	272	conserved hypothetical protein
SAG1300	57	conserved hypothetical protein
SAG1301	121	ribosomal protein L7/L12
SAG1302	166	ribosomal protein L10
SAG1303	702	ATP-dependent Clp protease, ATP-binding subunit
SAG1304	32	hypothetical protein
SAG1305	314	homocysteine S-methyltransferase MmuM, putative
SAG1306	458	amino acid permease
SAG1307	216	hypothetical protein
SAG1308	167	hypothetical protein
SAG1309	30	hypothetical protein
SAG1310	182	transcriptional regulator, TetR family
SAG1311	198	GTP-binding protein
SAG1312	408	ATP-dependent Clp protease, ATP-binding subunit ClpX
SAG1313	56	conserved hypothetical protein
SAG1314	164	dihydrofolate reductase
SAG1315	279	thymidylate synthase
SAG1316	390	HMG-CoA synthase
SAG1317	427	3-hydroxy-3-methylglutaryl-CoA reductase
SAG1318	149	conserved hypothetical protein
SAG1319	214	hemolysin III, putative
SAG1320	304	conserved hypothetical protein TIGR00147
SAG1321	284	glutathione S-transferase family protein, putative
SAG1322	72	conserved domain protein

ORF	Size	Annotati n
	(a.a.)	Augutau u
SAG1323	331	isopentenyl-diphosphate delta-isomerase
SAG1324	330	phosphomevalonate kinase
SAG1325	314	diphosphomevalonate decarboxylase
SAG1326	292	mevalonate kinase, putative
SAG1327	409	sensor histidine kinase
SAG1328	228	DNA-binding response regulator
SAG1329	208	GTP pyrophosphokinase family protein
SAG1330	68	hypothetical protein
SAG1331	979	R5 protein
SAG1332	146	transcriptional regulator, MarR family, putative
SAG1333	690	5'-nucleotidase family protein
SAG1334	136	polypeptide deformylase, putative
SAG1335	449	NADP-specific glutamate dehydrogenase
SAG1336	169	membrane protein, putative
SAG1337	589	ABC transporter, ATP-binding/permease protein
SAG1338	579	ABC transporter, ATP-binding/permease protein
SAG1339	157	acetyltransferase, GNAT family
SAG1340	622	ABC transporter, ATP-binding protein
SAG1341	402	polyA polymerase family protein
SAG1342	282	DegV family protein
SAG1343	126	protein of unknown function
SAG1344	177	hypothetical protein
SAG1345	164	conserved hypothetical protein
SAG1346	654	PTS system, fructose specific IIABC components
SAG1347	303	1-phosphofructokinase
SAG1348	247	lactose phosphotransferase system repressor
SAG1349	411	beta-lactam resistance factor
SAG1350	544	surface antigen-related protein
SAG1351	307	2-dehydropantoate 2-reductase, putative
SAG1352	356	regulatory protein, putative
SAG1353	330	pyridine nucleotide-disulphide oxidoreductase family protein
SAG1354	251	tRNA (guanine-N1)-methyltransferase
SAG1355	172	16S rRNA processing protein RimM
SAG1356	503	transcriptional regulator, RofA family
SAG1357	80	KH domain protein
SAG1358	90	ribosomal protein S16
SAG1359	415	permease, putative
SAG1360	236	ABC transporter, ATP-binding protein
SAG1361	414	conserved hypothetical protein
SAG1362	532	carbamoyl-phosphate synthase, large subunit, putative
SAG1363	356	carbamoyl-phosphate synthase, small subunit
SAG1364	173	pyrimidine operon regulatory protein
SAG1365	296	ribosomal large subunit pseudouridine synthase, RluD subfamily
SAG1366	154	lipoprotein signal peptidase
SAG1367	301	transcriptional regulator, LysR family
SAG1368	94	ribosomal protein L27
SAG1369	112	conserved hypothetical protein
SAG1370	104	ribosomal protein L21



OPE	l c:	
ORF	Size	Annotati n
SAG1371	(a.a.)	
SAG1371 SAG1372	392 404	conserved hypothetical protein
SAG1372 SAG1373	381	thiamine biosynthesis protein ThiI
		cysteine desulphurase
SAG1374	150	conserved hypothetical protein
SAG1375	449	glutathione reductase
SAG1376	111	conserved hypothetical protein
SAG1377	388	chorismate synthase
SAG1378	355	3-dehydroquinate synthase
SAG1379	225	3-dehydroquinate dehydratase
SAG1380	385	conserved hypothetical protein
SAG1381	714	sulfatase
SAG1382	119	ribosomal protein L20
SAG1383	66	ribosomal protein L35
SAG1384	176	translation initiation factor IF-3
SAG1385	227	cytidylate kinase
SAG1386	174	conserved hypothetical protein
SAG1387	65	ferredoxin, 4Fe-4S
SAG1388	163	conserved hypothetical protein
SAG1389	406	peptidase T
SAG1390	544	polysaccharide biosynthesis protein, putative
SAG1391	484	UDP-N-acetylmuramoylalanyl-D-glutamate2,6-diaminopimelate
		ligase
SAG1392	264	iron compound ABC transporter, ATP-binding protein
SAG1393	310	iron compound ABC transporter, substrate-binding protein
SAG1394	341	iron compound ABC transporter, permease protein
SAG1395	333	iron compound ABC transporter, permease protein
SAG1396	217	conserved hypothetical protein
SAG1397	311	inorganic pyrophosphatase, manganese-dependent
SAG1398	262	pyruvate formate-lyase-activating enzyme
SAG1399	444	CBS domain protein
SAG1400	188	conserved hypothetical protein
SAG1401	311	conserved hypothetical protein TIGR01212
SAG1402	213	PAP2 family protein
SAG1403	194	membrane protein, putative
SAG1404	308	cell wall surface anchor family protein
SAG1405	294	sortase family protein
SAG1406	293	sortase family protein
SAG1407	705	cell wall surface anchor family protein
SAG1408	901	cell wall surface anchor family protein
SAG1409	NA	rogB protein, authentic frameshift
SAG1410	379	glycosyl transferase, group 1 family protein
SAG1411	282	glycosyl transferase, group 2 family protein
SAG1412	474	polysaccharide biosynthesis protein
SAG1413	454	membrane protein, putative
SAG1414	308	glycosyl transferase, group 2 family protein
SAG1415	311	glycosyl transferase, group 2 family protein
SAG1416	352	nucleotide sugar dehydratase, putative
SAG1417	240	nucleotidyl transferase, putative



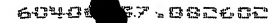
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ORF	Size (a.a.)	Ann tation
SAG1418	274	polysaccharide biosynthesis protein, putative
SAG1419	577	lipoprotein, putative
SAG1420	117	conserved hypothetical protein
SAG1421	243	glycosyl transferase, group 2 family protein
SAG1422	313	glycosyl transferase, group 2 family protein
SAG1423	384	glycosyl transferase, putative
SAG1424	284	dTDP-4-dehydrorhamnose reductase
SAG1425	113	conserved hypothetical protein
SAG1426	369	1 2 2 2 2 2 2 2 2 2 2 2 2 2 2 2 2 2 2 2
SAG1427	602	DNA primase
SAG1428	125	large conductance mechanosensitive channel protein
SAG1429	58	ribosomal protein S21
SAG1430	167	conserved hypothetical protein
SAG1431	268	amino acid ABC transporter, amino acid-binding protein
SAG1432	347	ammonium transporter family protein
SAG1433	375	conserved hypothetical protein
SAG1434	328	rhodanese family protein
SAG1435	101	conserved hypothetical protein
SAG1436	457	glycerol-3-phosphate transporter, putative
SAG1437	55	hypothetical protein
SAG1438	754	glycogen phosphorylase
SAG1439	498	4-alpha-glucanotransferase
SAG1440	342	maltose operon repressor MalR, putative
SAG1441	415	maltose/maltodextrin ABC transporter, maltose/maltodextrin-
	ļ	binding protein
SAG1442	456	maltose ABC transporter, permease protein
SAG1443	278	maltose ABC transporter, permease protein
SAG1444	490	proton/peptide symporter family protein
SAG1445	NA	MutT/nudix family protein, authentic frameshift
SAG1446	62	hypothetical protein
SAG1447	441	conserved hypothetical protein
SAG1448	502	glycosyl transferase, group 1 family protein
SAG1449	795	preprotein translocase SecA subunit, putative
SAG1450	330	
SAG1451	494	
SAG1452	514	
SAG1453	409	1-AA
SAG1454	398	<u> </u>
SAG1455	295	glycosyl transferase, group 2 family protein
SAG1456	NA	
SAG1457	129	IS1381, transposase OrfB
SAG1458	127	IS1381, transposase OrfA
SAG1459	413	glycosyl transferase family 8
SAG1460	401	glycosyl transferase, family 8
SAG1461	335	conserved hypothetical protein
SAG1462	970	
SAG1463	NA	transcriptional regulator, RofA family, authentic point mutation
SAG1464	663	excinuclease ABC, B subunit

ORF	Size	Annotation
	(a.a.)	THE PROPERTY OF THE PROPERTY O
SAG1465	306	protease, putative
SAG1466	727	glutamine ABC transporter, glutamine-binding protein/permease
		protein
SAG1467	246	glutamine ABC transporter, ATP-binding protein, GlnQ putative
SAG1468	116	conserved hypothetical protein
SAG1469	52	conserved hypothetical protein
SAG1470	437	GTP-binding protein, GTP1/Obg family
SAG1471	42	conserved hypothetical protein
SAG1472	413	aminopeptidase PepS
SAG1473	192	cell wall surface anchor family protein
SAG1474	680	amidase family protein
SAG1475	240	ribosomal small subunit pseudouridine synthase A
SAG1476	280	oxidoreductase, aldo/keto reductase family
SAG1477	224	nitroreductase family protein
SAG1478	130	lactoylglutathione lyase
SAG1479	308	glycosyl transferase, group 2 family protein
SAG1480	462	amino acid permease
SAG1481	155	SsrA-binding protein
SAG1482	801	exoribonuclease, VacB/Rnb family
SAG1483	78	preprotein translocase, SecG subunit
SAG1484	48	ribosomal protein L33
SAG1485	389	multi-drug resistance protein
SAG1486	548	membrane protein, putative
SAG1487	233	ABC transporter, ATP binding protein
SAG1488	195	dephospho-CoA kinase
SAG1489	273	formamidopyrimidine-DNA glycosylase
SAG1490	282	transcriptional regulator, MutR family
SAG1491	530	hypothetical protein
SAG1492	58	hypothetical protein
SAG1493	66	hypothetical protein
SAG1494	32	<del>                                     </del>
SAG1495	81	CAAX amino terminal protease family protein
SAG1496	110	hypothetical protein
SAG1497	37	
SAG1498	133	
SAG1499	299	<u> </u>
SAG1500	132	diacylglycerol kinase
SAG1501	161	conserved hypothetical protein TIGR00043
SAG1502	268	tetracenomycin polyketide synthesis O-methyltransferase TcmP,
		putative
SAG1503	39	hypothetical protein
SAG1504	38	+ <del></del>
SAG1505	158	+
SAG1506	267	hypothetical protein
SAG1507	345	<del> </del>
SAG1508	590	
	<u> </u>	
SAG1509	71	conserved hypothetical protein

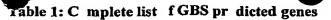
ORF	Size	Annotation
ORF		Annotation
SAG1511	(a.a.) 284	consequed hymothetical protein
SAG1511	185	conserved hypothetical protein ribosome recycling factor
SAG1513	242	uridylate kinase
SAG1514	226	peptide ABC transporter, ATP-binding protein
SAG1515	262	peptide ABC transporter, ATP-binding protein
SAG1516	255	peptide ABC transporter, permease protein
SAG1517	314	peptide ABC transporter, permease protein
SAG1518	538	peptide ABC transporter, peptide-binding protein
SAG1519	229	ribosomal protein L1
SAG1520	141	ribosomal protein L11
SAG1521	388	transposase, IS30 family, putative
SAG1522	460	transporter, major facilitator family
SAG1523	404	peptidase, M20/M25/M40 family
SAG1524	294	transcriptional regulator, LysR family
SAG1525	117	conserved hypothetical protein
SAG1526	178	IS861, transposase OrfA
SAG1527	277	IS861, transposase OrfB
SAG1528	571	chorismate binding enzyme .
SAG1529	816	
SAG1530	267	peptidyl-prolyl cis-trans isomerase, cyclophilin-type
SAG1531	277	manganese ABC transporter, permease protein
SAG1532	.238	manganese ABC transporter, ATP-binding protein
SAG1533	308	manganese ABC transporter, manganese-binding adhesion
		liprotein
SAG1534	215	iron-dependent transcriptional regulator
SAG1535	229	5-methylthioadenosine nucleosidase/S-adenosylhomocysteine
		nucleosidase
SAG1536	89	conserved hypothetical protein
SAG1537	184	MutT/nudix family protein
SAG1538	459	UDP-N-acetylglucosamine pyrophosphorylase
SAG1539	31	hypothetical protein
SAG1540	137	conserved hypothetical protein
SAG1541	125	glyoxalase family protein
SAG1542	318	oxidoreductase, Gfo/Idh/MocA family
SAG1543	NA	conserved hypothetical protein, authentic frameshift
SAG1544	232	gluconate 5-dehydrogenase, putative
SAG1545	78	conserved hypothetical protein
SAG1546	82	conserved hypothetical protein
SAG1547	166	
SAG1548	422	glycosyl transferase, group 2 family protein
SAG1549	127	
SAG1550	129	
SAG1551	67	hypothetical protein
SAG1552	719	conserved hypothetical protein
SAG1553	477	hypothetical protein
SAG1554	225	hypothetical protein
SAG1555	231	hypothetical protein
SAG1556	445	
חרכזטעם	L 443	branched-chain amino acid transport system II carrier protein



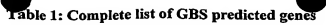
ORF	Size	Ann tation
OKF	(a.a.)	Ann tanon
SAG1557	665	methionyl-tRNA synthetase
SAG1558	291	tellurite resistance protein TehB
SAG1559	231	membrane protein, putative
SAG1560	40	hypothetical protein
SAG1561	405	PTS system, IIC component, putative
SAG1562	280	conserved hypothetical protein
SAG1563	275	exodeoxyribonuclease
SAG1564	118	conserved hypothetical protein
SAG1565	158	methylated-DNAprotein-cysteine S-methyltransferase
SAG1566	393	D-isomer specific 2-hydroxyacid dehydrogenase family protein
SAG1567	182	acetyltransferase, GNAT family
SAG1568	NA 211	phosphoserine aminotransferase, authentic frameshift
SAG1569 SAG1570	211	copper homeostasis protein CutC, putative
	53	conserved hypothetical protein
SAG1571 SAG1572	287	hypothetical protein
		tetrapyrrole methylase family protein
SAG1573	108 287	conserved hypothetical protein
SAG1574		DNA polymerase III, delta prime subunit, putative
SAG1575	211	thymidylate kinase
SAG1576	267	transposase, IS30 family, putative, truncation
SAG1577	219	AcuB family protein
SAG1578	236	branched-chain amino acid ABC transporter, ATP-binding protein
SAG1579	254	branched-chain amino acid ABC transporter, ATP-binding protein
SAG1580	317	branched-chain amino acid ABC transporter, permease protein.
SAG1581 SAG1582	289	branched-chain amino acid ABC transporter, permease protein
SAG1362	388	branched-chain amino acid ABC transporter, amino acid-binding protein
SAG1583	81	conserved hypothetical protein
SAG1584	377	
SAG1585	196	· <del></del>
SAG1586	209	
SAG1587	389	aminotransferase, class I
SAG1588	182	RNA methyltransferase, TrmH family, group 2
SAG1589	450	
SAG1589	449	<del></del>
SAG1591	475	cation uptake protein, Trk family
SAG1591	83	conserved hypothetical protein TIGR00278
SAG1593	240	
SAG1593	194	
SAG1595	235	<u> </u>
SAG1596	246	
SAG1590	157	
SAG1598	173	
SAG1599	324	<del></del>
SAG1600		
SAG1601	264	
SAG1601	190	<u></u>
	180	
SAG1603	173	transcriptional regulator, biotin repressor family



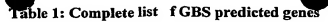
ORF	Siz	Annotation
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SAG1604	229	membrane protein, putative
SAG1605	167	conserved hypothetical protein
SAG1606	247	RNA methyltransferase, TrmH family
SAG1607	92	acylphosphatase
SAG1608	310	lipoprotein, putative
SAG1609.	221	amino acid ABC transporter, permease protein
SAG1610	285	amino acid ABC transporter, substrate-binding protein
SAG1611	486	amidase family protein
SAG1612	160	transcription elongation factor GreA
SAG1613	600	conserved hypothetical protein
SAG1614	167	acetyltransferase, GNAT family
SAG1615	443	UDP-N-acetylmuramatealanine ligase
SAG1616	205	conserved hypothetical protein
SAG1617	32	hypothetical protein
SAG1618	1032	Snf2 family protein
SAG1619	377	IS1548, transposase
SAG1620	436	
SAG1621	300	primosomal protein Dnal
SAG1622	391	conserved hypothetical protein
SAG1623	159	conserved hypothetical protein TIGR00244
SAG1624	501	sensor histidine kinase CsrS
SAG1625	229	DNA-binding response regulator CsrR
SAG1626	177	conserved hypothetical protein
SAG1627	296	heat shock protein HtpX
SAG1628	184	lemA protein
SAG1629	237	glucose-inhibited division protein B
SAG1630	459	sodium transport family protein
SAG1631	223	potassium uptake protein, Trk family, putative
SAG1632	276	cobalt transport family protein
SAG1633	558	ABC transporter, ATP-binding protein
SAG1634	212	conserved hypothetical protein
SAG1635	402	sodium:dicarboxylate symporter family protein
SAG1636	455	branched-chain amino acid transport system II carrier protein
SAG1637	351	alcohol dehydrogenase, zinc-containing
SAG1638	230	
SAG1639	356	· · · · · · · · · · · · · · · · · · ·
		· · · · · · · · · · · · · · · · · · ·
SAG1640	458	. <del></del>
SAG1641	274	
SAG1642	277	ABC transporter, substrate-binding protein
SAG1643	229	
SAG1644	37	hypothetical protein
SAG1645	238	
SAG1646	32	
SAG1647	328	<u> </u>
SAG1648	178	
SAG1649	37	
SAG1650	329	
SAG1651	192	dihydroxyacetone kinase family protein



0.77		
ORF	Size	Annotation
01010	(a.a.)	
SAG1652	124	conserved hypothetical protein
SAG1653	237	glycerol uptake facilitator protein
SAG1654	134	conserved hypothetical protein
SAG1655	237	transcriptional regulator, MerR family
SAG1656	369	conserved hypothetical protein
SAG1657	83	hypothetical protein
SAG1658	244	conserved hypothetical protein
SAG1659	118	iojap-related protein
SAG1660	173	isochorismatase family protein
SAG1661	195	conserved hypothetical protein TIGR00488
SAG1662	210	conserved hypothetical protein TIGR00482
SAG1663	105	conserved hypothetical protein TIGR00253
SAG1664	372	GTP-binding protein
SAG1665	177	hydrolase, haloacid dehalogenase-like family
SAG1666	304	membrane protein, putative
SAG1667	480	glutamyl-tRNA(Gln) amidotransferase, B subunit
SAG1668	488	glutamyl-tRNA(Gln) amidotransferase, A subunit
SAG1669	100	glutamyl-tRNA(Gln) amidotransferase, C subunit
SAG1670	881	pyruvate phosphate dikinase
SAG1671	276	protein of unknown function -
SAG1672	170	CBS domain protein
SAG1673	321	3-hydroxyacyl-CoA dehydrogenase family protein
SAG1674	182	isochorismatase family protein
SAG1675	261	transcriptional regulator CodY, putative
SAG1676	403	aminotransferase, class I
SAG1677	150	conserved hypothetical protein
SAG1678	460	hydrolase, haloacid dehalogenase-like family
SAG1679	320	asparaginase family protein
SAG1680	292	
SAG1681	304	
SAG1682	671	
SAG1683	512	
SAG1684	366	<u> </u>
SAG1685	119	
SAG1686	335	
SAG1687	842	<u></u>
SAG1688	315	
SAG1689	293	
SAG1690	639	· <del></del>
SAG1691	479	
SAG1692	320	
SAG1693	144	
SAG1694	129	
SAG1695		
	186	
SAG1696	38	
SAG1697	48	
SAG1698	99	
SAG1699	30	hypothetical protein



ORF	Size (a.a.)	Annotation
SAG1700	76	hypothetical protein
SAG1701	56	hypothetical protein
SAG1702	41	hypothetical protein
SAG1703	54	hypothetical protein
SAG1704	150	cytidine/deoxycytidylate deaminase family protein
SAG1705	NA	peptidase, M24 family, authentic point mutation
SAG1706	238	conserved hypothetical protein
SAG1707	499	drug resistance transporter, EmrB/QacA family
SAG1708	38	hypothetical protein
SAG1709	942	excinuclease ABC, A subunit
SAG1710	223	conserved hypothetical protein
SAG1711	314	magnesium transporter, CorA family
SAG1712	79	ribosomal protein S18
SAG1713	163	single-strand binding protein
SAG1714	95	ribosomal protein S6
SAG1715	374	A/G-specific adenine glycosylase
SAG1716	197	transcriptional regulator, Cro/CI family
SAG1717	104	thioredoxin
SAG1718	166	PAP2 family protein
SAG1719	779	MutS2 family protein
SAG1720	180	conserved hypothetical protein
SAG1721	103	conserved hypothetical protein
SAG1722	297	ribonuclease HIII
SAG1723	197	signal peptidase I
SAG1724	806	helicase, putative
SAG1725	160	conserved hypothetical protein
SAG1726	364	DNA-damage-inducible protein P
SAG1727	770.	formate acetyltransferase
SAG1728	124	FMN-binding protein
SAG1729	309	conserved hypothetical protein
SAG1730	251	conserved hypothetical protein
SAG1731	298	membrane protein, putative
SAG1732	282	glycerol uptake facilitator protein, putative
SAG1733	150	universal stress protein family
SAG1734	400	transporter, putative
SAG1735	219	transcriptional regulator, Crp/Fnr family
SAG1736	761	
SAG1737	119	
SAG1738	326	polyprenyl synthetase family protein
SAG1739	582	
SAG1740	572	
SAG1741	339	<u> </u>
SAG1742	475	cytochrome d oxidase, subunit I
SAG1743	402	· · · · · · · · · · · · · · · · · · ·
SAG1744	299	
SAG1745	148	
SAG1746	35	
SAG1747	99	



ORF	Size	Ann tati n
	(a.a.)	
SAG1748	396	cyclopropane-fatty-acyl-phospholipid synthase
SAG1749	241	transcriptional regulator, merR family
SAG1750	195	exonuclease
SAG1751	178	conserved hypothetical protein
SAG1752	390	conserved hypothetical protein TIGR00275
SAG1753	260	conserved hypothetical protein
SAG1754	89	ribosomal protein S14
SAG1755	38	hypothetical protein
SAG1756	341	conserved hypothetical protein
SAG1757	336	O-sialoglycoprotein endopeptidase family protein
SAG1758	135	ribosomal-protein-alanine acetyltransferase, putative
SAG1759	230	protein of unknown function
SAG1760	76	conserved hypothetical protein
SAG1761	559	metallo-beta-lactamase superfamily protein
SAG1762	169	conserved hypothetical protein
SAG1763	448	glutamine synthetase, type I
SAG1764	123	transcriptional regulator GlnR
SAG1765	179	conserved hypothetical protein
SAG1766	398	phosphoglycerate kinase
SAG1767	289	acid phosphatase
SAG1768	336	glyceraldehyde 3-phosphate dehydrogenase
SAG1769	692	translation elongation factor G
SAG1770	156	ribosomal protein S7
SAG1771	137	ribosomal protein S12
SAG1772	270	pur operon repressor
SAG1773	313	HD domain protein
SAG1774	424	conserved hypothetical protein
SAG1775	210	conserved hypothetical protein
SAG1776	220	ribulose-phosphate 3-epimerase
SAG1777	290	conserved hypothetical protein TIGR00157
SAG1778	283	rRNA (guanine-N1-)-methyltransferase, putative
SAG1779	290	dimethyladenosine transferase
SAG1780	163	
SAG1781	186	+
SAG1782	260	
SAG1783	90	
SAG1784	130	
SAG1785	430	
SAG1786	130	
SAG1787	420	
SAG1788	79	D-alanyl carrier protein
SAG1789	421	<u> </u>
SAG1790	511	· · · · · · · · · · · · · · · · · · ·
SAG1791	395	
SAG1792	224	
SAG1793	44	
SAG1794		
SAG1795	388	transposase, IS30 family, putative

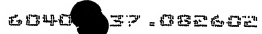


Sizo	Annotation
- 1	Annotation .
	amino acid ABC transporter, permease protein
	amino acid ABC transporter, ATP-binding protein
	hypothetical protein
	xylulose-5-phosphate/fructose-6-phosphate phosphoketolase
	conserved hypothetical protein
	transcriptional antiterminator, BglG family
	conserved hypothetical protein
	carbohydrate kinase, FGGY family
	hypothetical protein
	PTS system, IIC component, putative
	glyoxylate reductase, NADH-dependent
	hypothetical protein
	sugar binding transcriptional regulator, LacI family
	transaldolase family protein
	carbohydrate isomerase, AraD/FucA family hexulose-6-phosphate isomerase, putative
<del></del>	hexulose-6-phosphate synthase, putative
	PTS system, IIA component
	PTS system, IIB component
	transport protein SgaT, putative
	hypothetical protein
	hypothetical protein
<del></del>	adenylosuccinate synthetase
	perfringolysin O regulator protein
	conserved hypothetical protein
	glutamatecysteine ligase/amino acid ligase, putative
	protein of unknown function
	protein of unknown function
+	chaperonin, 33 kDa
	NifR3/Smm1 family protein
<del></del>	deoxynucleoside kinase family protein
	phosphinothricin N-acetyltransferase
	. <b> </b>
	<u> </u>
<del></del>	<u></u>
1224	
	· ····································
	Size         (a.a.)           575         407           39         792           363         559           253         505           329         483           318         339           327         215           238         287           221         161           92         479           205         157           430         340           224         750           272         418           291         325           213         163           815         154           153         346           256         186           510         134           61         468           109         136           112         76           124         124

ORF	Size	Annotation
OKF	(a.a.)	Autotation
SAG1844	911	conserved hypothetical protein
SAG1845	42	hypothetical protein
SAG1846	158	hypothetical protein
SAG1847	227	conserved hypothetical protein
SAG1848	114	conserved hypothetical protein
SAG1849	115	hypothetical protein
SAG1850	101	hypothetical protein
SAG1851	111	conserved domain protein
SAG1852	420	conserved domain protein
SAG1853	180	prophage LambdaSa2, protease, putative
SAG1854	380	conserved hypothetical protein
SAG1855	570	prophage LambdaSa2, terminase large subunit, putative
SAG1856	161	hypothetical protein
SAG1857	119	prophage LambdaSa2, HNH endonuclease family protein
SAG1858	95	
SAG1859	180	hypothetical protein prophage LambdaSa2, site-specific recombinase, phage integrase
3AG1639	160	family
SAG1860	154	conserved hypothetical protein
SAG1861	119	prophage LambdaSa2, transcriptional regulator, Cro/CI family
SAG1862	86	hypothetical protein
SAG1863	138	prophage LambdaSa2, single-strand binding protein
SAG1864	68	hypothetical protein
SAG1865	74	conserved hypothetical protein
SAG1866	109	conserved hypothetical protein
SAG1867	163	conserved hypothetical protein
SAG1868	134	hypothetical protein
SAG1869	437	prophage LambdaSa2, type II DNA modification
	<u> </u>	methyltransferase, putative
SAG1870	273	prophage LambdaSa2, DNA replication protein DnaC, putative
SAG1871	248	prophage LambdaSa2, bacteriophage replication protein/hypothetical protein, truncation/fusion
SAG1872	200	
SAG1873	1	
SAG1874	443 87	prophage LambdaSa2, replicative DNA helicase hypothetical protein
SAG1875	94	<u> </u>
SAG1876 SAG1877	176 236	· [-^
SAG1877 SAG1878		
<u> </u>	102	
SAG1879 SAG1880	156	<u>,  </u>
<del></del>	54	
SAG1881 SAG1882	51	
	120	<u> </u>
SAG1883	128	
SAG1884	134	
SAG1885	356	prophage LambdaSa2, site-specific recombinase, phage integrase family
SAG1886	32	. <u> </u>
SAG1887	689	



ORF	Size	Annotati n
024	(a.a.)	
SAG1888	78	hypothetical protein
SAG1889	317	microcin immunity protein MccF, putative
SAG1890	631	endopeptidase O
SAG1891	327	oxidoreductase, Gfo/Idh/MocA family
SAG1892	358	membrane protein, putative
SAG1893	59	hypothetical protein
SAG1894	214	cyclic nucleotide-binding domain protein
SAG1895	204	polypeptide deformylase
SAG1896	333	sugar binding transcriptional regulator RegR
SAG1897	634	conserved hypothetical protein
SAG1898	271	PTS system, IID component
SAG1899		PTS system, IIC component
SAG1900	164	
SAG1901	398	glucuronyl hydrolase
SAG1902	144	PTS system, IIA component
SAG1903	34	hypothetical protein
SAG1904	270	oxidoreductase, short-chain dehydrogenase/reductase family
SAG1905	212	conserved hypothetical protein
SAG1906	335	carbohydrate kinase, PfkB family
SAG1907	212	2-dehydro-3-deoxyphosphogluconate aldolase/4-hydroxy-2-
04.01000	400	oxoglutarate aldolase
SAG1908	499	hypothetical protein
SAG1909	204	<u> </u>
SAG1910	141	transcriptional regulator, MarR family
SAG1911	1468	DNA polymerase III, alpha subunit, Gram-positive type
SAG1912	194	N-acetylmuramoyl-L-alanine amidase, family 4 protein
SAG1913	617	prolyl-tRNA synthetase
SAG1914	419	membrane-associated zinc metalloprotease, putative
SAG1915	264	phosphatidate cytidylyltransferase
SAG1916	250	
SAG1917	113	preprotein translocase, YajC subunit
SAG1918	114	
SAG1919	387	malate oxidoreductase
SAG1920	445	
SAG1921	508	
SAG1922	229	
SAG1923	331	
SAG1924	535	<u></u>
SAG1925	377	
SAG1926	283	<u></u>
SAG1927	298	
SAG1928	325	
SAG1929	310	
SAG1930	171	
SAG1931	141	
SAG1932	816	
SAG1933	482	
SAG1934	101	PTS system, IIB component, putative



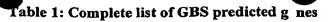
ORF	Size	Annotation
Old	(a.a.)	Amilotation
SAG1935	157	PTS system, IIA component, putative
SAG1936	258	lactose phosphotransferase system repressor
SAG1937	NA	streptococcal histidine triad family protein, degenerate
SAG1938	307	adhesion lipoprotein
SAG1939	147	protein of unknown function TIGR00256
SAG1940	738	GTP pyrophosphokinase family protein
SAG1941	800	2`,3`-cyclic-nucleotide 2`-phosphodiesterase
SAG1942	151	nrdI protein
SAG1943	345	conserved hypothetical protein
SAG1944	165	conserved hypothetical protein
SAG1945	345	iron ABC transporter, iron-binding protein
SAG1946	257	DNA-binding response regulator
SAG1947	549	conserved hypothetical protein
SAG1948	275	PTS system, IID component
SAG1949	269	PTS system, IIC component
SAG1950	163	PTS system, IIB component
SAG1951	141	PTS system, IIA component, putative
SAG1952	353	membrane protein, putative
SAG1953	60	hypothetical protein
SAG1954	384	membrane protein, putative
SAG1955	282	ABC transporter, ATP-binding protein
SAG1956	96	conserved hypothetical protein, truncation
SAG1957	250	response regulator
SAG1958	276	conserved hypothetical protein
SAG1959	727	PTS system, IIABC components
SAG1960	551	sensor histidine kinase
SAG1961	225	phosphate regulon response regulator PhoB
SAG1962	218	phosphate transport system regulatory protein PhoU, putative
SAG1963	253	phosphate ABC transporter, ATP-binding protein
SAG1964	292	phosphate ABC transporter, permease protein
SAG1965	281	phosphate ABC transporter, permease protein
SAG1966	293	hemolysin precursor, putative
SAG1967	195	hypothetical protein
SAG1968	• 246	
SAG1969	317	· · · · · · · · · · · · · · · · · · ·
SAG1970	102	
SAG1971	41	hypothetical protein
SAG1972	238	- <del></del>
SAG1973	156	· <del>                                    </del>
SAG1974	152	
SAG1975	47	
SAG1976	156	· <del>  · · · · · · · · · · · · · · · · · ·</del>
SAG1977	163	- <del></del>
SAG1978	422	
SAG1979	253	
SAG1980	300	
SAG1981	68	
SAG1982	359	
DAU1702	779	Lanscriptional regulator, Cro/Cr failing

# Table 1: C mplete list of GBS predicted genes

ORF	Size	Annotation
G 1 G 1 G G	(a.a.)	
SAG1983	105	conserved hypothetical protein
SAG1984	188	conserved hypothetical protein TIGR00730
SAG1985	51	hypothetical protein
SAG1986	375	site-specific recombinase, phage integrase family
SAG1987	61	conserved hypothetical protein
SAG1988	342	conserved hypothetical protein
SAG1989	139	hypothetical protein .
SAG1990	127	hypothetical protein
SAG1991	204	transcriptional regulator, Cro/CI family
SAG1992	518	protein of unknown function
SAG1993	373	site-specific recombinase, phage integrase family
SAG1994	108	conserved hypothetical protein
SAG1995	210	hypothetical protein
SAG1996	263	cell wall surface anchor family protein, putative
SAG1997	182	hypothetical protein
SAG1998	457	hypothetical protein
SAG1999	47	hypothetical protein
SAG2000	666	membrane protein, putative
SAG2001	756	conjugal transfer protein, interruption-C
SAG2002	129	IS1381, transposase OrfB
SAG2003	127	IS1381, transposase OrfA
SAG2004	67	conjugal transfer protein, interruption-N
SAG2005	136	conserved hypothetical protein
SAG2006	88	conserved hypothetical protein
SAG2007	317	conserved hypothetical protein
SAG2008	84	conserved hypothetical protein
SAG2009	88	conserved hypothetical protein
SAG2010	157	hypothetical protein
SAG2011	160	conserved hypothetical protein
SAG2012	90	hypothetical protein
SAG2013	189	·
SAG2014	449	
SAG2015	99	transcriptional regulator, Cro/CI family
SAG2016	125	
SAG2017	429	
SAG2018	553	
SAG2019	153	-V
SAG2020	98	
SAG2021	826	
SAG2022	417	
SAG2023	546	
SAG2024	130	· · · · · · · · · · · · · · · · ·
SAG2025	522	
SAG2026	240	
SAG2027	205	
SAG2028	36	
SAG2029	284	
SAG2030	130	
L 3A 32030	1 130	I Hypometical protetti

# Table 1: Complete list of GBS predicted genes

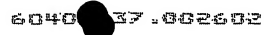
022	C'	A							
ORF	Size (a.a.)	Annotati n							
SAG2031	202	hypothetical protein							
SAG2032	111	conserved hypothetical protein							
SAG2033	162	acetyltransferase, GNAT family							
SAG2034	247	membrane protein, putative							
SAG2035	300	ABC transporter, ATP-binding protein							
SAG2036	68	hypothetical protein							
SAG2037	358	transcriptional regulator, Cro/CI family							
SAG2038	204								
SAG2039	98	conserved hypothetical protein							
SAG2040	186	conserved hypothetical protein TIGR00730							
SAG2041	287	protease, putative							
SAG2042	100	rhodanese family protein							
SAG2043	255	cAMP factor							
SAG2044	62	hypothetical protein							
SAG2045	179	<del>                                     </del>							
SAG2045	361	glycerol dehydrogenase, putative							
SAG2047	235	conserved hypothetical protein							
SAG2047 SAG2048	614	5-methyltetrahydrofolatehomocysteine methyltransferase,							
SAG2048		putative							
SAG2049	745	5-methyltetrahydropteroyltriglutamatehomocysteine methyltransferase							
SAG2050	107								
SAG2051	230								
SAG2052	41	hypothetical protein							
SAG2053	1570	serine protease, subtilase family, putative							
SAG2054	228								
SAG2055	462	sensor histidine kinase							
SAG2056	202								
SAG2057	833								
SAG2058	415								
SAG2059	281								
SAG2060	398	<u> </u>							
SAG2061	401	glycosyl transferase, family 8							
SAG2062	179	<u> </u>							
SAG2063	630								
SAG2064	57								
SAG2065	50								
SAG2066	773								
SAG2067	294								
SAG2068	546								
SAG2069	403								
SAG2009									
<del></del>	223								
SAG2071	400								
SAG2072	259								
SAG2073	245								
SAG2074	540								
SAG2075	94								
SAG2076	267	ABC transporter, ATP-binding protein							



ODE	Q:	Ann Action								
ORF	Size	Ann tation								
G A C2077	(a.a.)	ADC								
SAG2077	298	ABC transporter, permease protein								
SAG2078	320	protein of unknown function/lipoprotein, putative								
SAG2079	265	hydrolase, haloacid dehalogenase-like family								
SAG2080	286	glyoxalase family protein								
SAG2081	243	conserved hypothetical protein								
SAG2082	205	anaerobic ribonucleoside-triphosphate reductase activating protein								
SAG2083	163	acetyltransferase, GNAT family								
SAG2084	310	virulence factor MviM, putative								
SAG2085	47	conserved hypothetical protein								
SAG2086	723	anaerobic ribonucleoside-triphosphate reductase								
SAG2087	495	membrane protein, putative								
SAG2088	40	hypothetical protein								
SAG2089	105	conserved hypothetical protein								
SAG2090	136	conserved hypothetical protein TIGR00250								
SAG2091	88	conserved hypothetical protein								
SAG2092	132	conserved hypothetical protein								
SAG2093	379	recA protein								
SAG2094	NA	competence/damage-inducible protein CinA, authentic frameshift								
SAG2095	183	DNA-3-methyladenine glycosylase I								
SAG2096	196	Holliday junction DNA helicase RuvA								
SAG2097	418	transporter, putative								
SAG2098	659	DNA mismatch repair protein HexB								
SAG2099	33	hypothetical protein								
SAG2100	67	cold shock protein, CSD family								
SAG2101	858	DNA mismatch repair protein HexA								
SAG2102	145	arginine repressor ArgR, putative								
SAG2103	563	arginyl-tRNA synthetase								
SAG2104	102	conserved hypothetical protein								
SAG2105	290	conserved hypothetical protein								
SAG2106	314	conserved hypothetical protein								
SAG2107	583	aspartyl-tRNA synthetase								
SAG2108	426	histidyl-tRNA synthetase								
SAG2109	60	ribosomal protein L32								
SAG2110	49									
SAG2111	173	conserved hypothetical protein								
SAG2112	494	site-specific recombinase, phage integrase family								
SAG2113	82	conserved hypothetical protein								
SAG2114	342	<u> </u>								
SAG2115	143									
SAG2116	151	conserved hypothetical protein								
SAG2117	71	hypothetical protein								
SAG2118	306									
SAG2118 SAG2119										
	373									
SAG2120	269									
SAG2121	223									
SAG2122	223	<u> </u>								
SAG2123	454	<u> </u>								
SAG2124	517	membrane protein, putative								

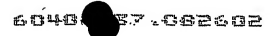
# Table 1: Complete list of GBS predicted genes

	· .								
ORF	Size	Annotati n							
	(a.a.)								
SAG2125	308	carbamate kinase							
SAG2126	332	omithine carbamoyltransferase							
SAG2127	431	sensor histidine kinase							
SAG2128	277	response regulator							
SAG2129	240	amino acid ABC transporter, ATP-binding protein							
SAG2130	504	amino acid ABC transporter, amino acid-binding protein/permea							
		protein							
SAG2131	847	membrane protein, putative							
SAG2132	247	conserved hypothetical protein							
SAG2133	118	conserved hypothetical protein							
SAG2134	772	membrane protein, putative							
SAG2135	179	transcriptional regulator, TetR family, putative							
SAG2136	98	conserved hypothetical protein							
SAG2137	203	ribosomal protein S4							
SAG2138	95	conserved hypothetical protein							
SAG2139	451	replicative DNA helicase							
SAG2140	150	ribosomal protein L9							
SAG2141	660	DHH family protein							
SAG2142	613	glucose inhibited division protein A							
SAG2143	203	membrane protein, putative							
SAG2144	373	tRNA (5-methylaminomethyl-2-thiouridylate)-methyltransfera							
SAG2145	222	L-serine dehydratase, iron-sulfur-dependent, beta subunit							
SAG2146	290	L-serine dehydratase, iron-sulfur-dependent, beta subunit							
SAG2147	234	protein of unknown function/lipoprotein, putative							
SAG2148	179	LysM domain protein							
SAG2149	264	cobalt transport family protein							
SAG2150	280	ABC transporter, ATP-binding protein							
SAG2151	279								
SAG2152	180	CDP-diacylglycerolglycerol-3-phosphate 3-							
51.102.02		phosphatidyltransferase							
SAG2153	427	peptidase, M16 family							
SAG2154	414	conserved hypothetical protein							
SAG2155	117	conserved hypothetical protein							
SAG2156	369	recF protein							
SAG2157	278	· · · · · · · · · · · · · · · · · · ·							
SAG2158	220								
SAG2159	493								
SAG2160	161	transcriptional regulator, ArgR family							
SAG2161	226								
SAG2162	234								
SAG2163	410								
SAG2164	136								
SAG2165	337								
SAG2166	475								
SAG2167	318								
SAG2167 SAG2168	341								
SAG2169		tryptophanyl-tRNA synthetase							
	230								
SAG2170	290	conserved hypothetical protein							

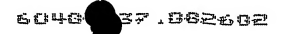


# Table 1: Complete list of GBS predicted genes

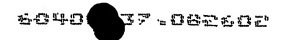
ORF	Size (a.a.)	Annotati n						
SAG2171	539	ABC transporter, ATP-binding protein						
SAG2172	859	ABC transporter, permease protein, putative						
SAG2173	159	conserved hypothetical protein TIGR00246						
SAG2174	409	serine protease						
SAG2175	257	partitioning protein, ParB family						



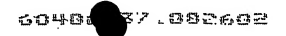
	Size	Signal	Sortase	Lipo-		Western		GBS	
ORF	(aa)	Peptide	motif	protein	Other	blot	FACS	specific	Annotation
SAG0017	447	+							pcsB
SAG0031	299	+	-						peptidase, M23/M37 family
SAG0032	434	+				+	+		group B streptococcal surface immunogenic protein
SAG0034	438	+		+		+	+		sugar ABC transporter, sugar-binding protein
SAG0051	126	+				+	+		MORN motif family protein
SAG0079	212				+	+	+		adenylate kmase
SAG0086	85			+				+	lipoprotein, putative
SAG0093	250	+				+	+		D-alanyl-D-alanine carboxypeptidase family protein
SAG0094	191	+							N-acetylmuramoyl-L-alanıne amıdase, famıly 4 protein
SAG0108	308	+							conserved hypothetical protein
									ribose ABC transporter, periplasmic D-ribose-binding
SAG0114	322	+		+					protein
SAG0124	356	+							sensor histidine kinase
SAG0132	294	+				+	+		SPFH domain/Band 7 family protein
SAG0134	96	+						+	hypothetical protein
SAG0146	395	+							penicullin-binding protein 4, putative
SAG0147	411	+							D-alanyl-D-alanine carboxypeptidase family protein
									oligopeptide ABC transporter, substrate-binding protein,
SAG0148	551	1	•	+		+	-		putative
SAG0166	123	+							conserved domain protein
SAG0176	94	+							conserved hypothetical protein
									oligopeptide ABC transporter, oligopeptide-binding
SAG0187	542	+	1	+		+	+		protein .
SAG0206	. 60	)		+				+	lipoprotein, putative
SAG0213	39	+						+	hypothetical protein
SAG0231	135	+							hypothetical protein
SAG0242	308	3		+		+	1 -		amino acid ABC transporter, amino acid-binding protein
SAG0245	152	2		+		+	·	+	protein of unknown function/lipoprotein, putative
SAG0255	31:	5 +							conserved hypothetical protein
SAG0257	5:	3		+			T	+	Inpoprotein, putative
SAG0265	23:	5 +				+	† -	+	conserved hypothetical protein
SAG0290	270	+			1	+	+		ABC transporter, substrate-binding protein
SAG0298	750	+	1		<b>†</b>	1			penicillin-binding protein 1A



	Size	Signal	Sortase	Lipo-		Western		GB\$	
ORF	(aa)	Peptide	motif	protein	Other	blot	FACS	specific	Annotation
SAG0306	535	+							KH domain protein
SAG0321	339	+							sensor histidine kinase, putative
SAG0329	106	+							PTS system, cellobiose-specific IIB component
SAG0368	435	+				+	+		protein of unknown function
SAG0371	167	+						+	hypothetical protein
SAG0383	334	+		+		+	-		protein of unknown function/lipoprotein, putative
SAG0392	521	+	+			+	+		cell wall surface anchor family protein
SAG0394	345				+				sensor histidine kinase
SAG0405	347	+		+		+	+		protein of unknown function/lipoprotein, putative
SAG0406	299	+							UTP-glucose-1-phosphate uridylyltransferase
SAG0407	338	+							glycerol-3-phosphate dehydrogenase (NAD(P)+)
SAG0416	1233	+	+			+	+		protease, putative
SAG0421	1055		+			+	-		cell wall surface anchor family protein
SAG0433	1389		+						surface protein Rib
SAG0437	123			+					lipoprotein, putative
SAG0451	149	+		+				+	bacteriocin transport accessory protein,putative
SAG0455	357	+							conserved hypothetical protein
SAG0472	126	+				+	-		rhodanese-like family protein
SAG0482	84	+							YGGT family protein
SAG0499	275				+				hemolysm A
SAG0503	279	+				+	+		lipase/acylhydrolase
SAG0504	200	+							conserved hypothetical protein
SAG0506	65	+						+	hypothetical protein
SAG0521	236	+							carboxymethylenebutenolidaso-related protein
SAG0535	506	+				+	+		zinc ABC transporter, zinc-binding adhesion liprotein
SAG0596	670				+				prophage LambdaSal, pblA protein, internal deletion
SAG0603	111				+				conserved hypothetical protein
SAG0604	239				+				prophage LambdaSal, lysin, putative
SAG0617	439				+				sensor histidine kinase VneS
SAG0624	574	+							septation ring formation regulator EzrA, putative
SAG0629	354	+							conserved domain protein
SAG0635	245	+				+			аси phosphatase, class В
SAG0638	109	+							cell wall surface anchor family protein, interruption-N

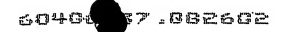


	Size	Signal	Sortase	Lipo-		Western		GBS	
ORF	(aa)	Peptide	motif	protein	Other	blot	FACS	specific	Annotation
SAG0645	554		+			+	+		cell wall surface anchor family protein
SAG0646	307	+	+			+	-		cell wall surface anchor family protein
SAG0647	305	+							sortase family protein
SAG0649	890		+			+	+		cell wall surface anchor family protein, putative
SAG0658	383	+		+					Irpoprotein, putative
SAG0675	171	+						•	putative secreted protein
SAG0676	885				+				proteinase, putative
SAG0677	1062		+						hypothetical protein
SAG0679	343	+		+		+	-		protein of unknown function
SAG0680	339	+				+	-		protein of unknown function
SAG0681	353	+							conserved domain protein
SAG0686	261	+				+	+		DNA-entry nuclease, putative
SAG0714	188	+						+	conserved hypothetical protein
SAG0717	266	+				+	+		amino acid ABC transporter, amino acid-binding protein
SAG0720	449				+				sensory box histidine kinase
SAG0738	132	+	,,						conserved hypothetical protein
SAG0739	143	+							conserved hypothetical protein
SAG0742	428				+	+	+		peptidase, U32 family
SAG0755	282	+							peptidase, U32 family
SAG0757	129	+		+		+	-	-	protein of unknown function/lipoprotein, putative
SAG0764	230				+	+	+		phosphoglycerate mutase family protein
SAG0765	681	+							penicillin-binding protein 2b
SAG0771	512	+	+			+	+	+	cell wall surface anchor family protein
SAG0776	276	+		+		<u> </u>			YaeC family protein, putative
SAG0777	528				+	+	+		ATP-dependent RNA helicase, DEAD/DEAH box family
SAG0785	330	+							conserved hypothetical protein
SAG0808	309	+		+		+	+		protease maturation protein, putative
SAG0824	417	+							polysaccharide deacetylase family protein
SAG0832	753	+				+	+		protein of unknown function
SAG0833	181	+				1		+ .	hypothetical protein
SAG0867	63	+							conserved hypothetical protein
SAG0868	285	+		<u> </u>		+	-		DNA-entry nuclease
SAG0886	319	+				+	+		protein of unknown function



	Size	Signal	Sortase	Lipo-		Western		GBS	
ORF	(aa)	Peptide	motif	protein	Other	blot	FACS	specific	Annotation
SAG0904	56	+						+	hypothetical protein
SAG0907	877	+		+		+	-		protein of unknown function/lipoprotein, putative
SAG0926	333	+							Tn916, NLP/P60 family protein
SAG0942	185	+				+	+		signal pepudase 1, putative
SAG0949	276	+		+		+	+		amino acid ABC transporter, amino acid-binding protein
SAG0954	349			+		+	-		protein of unknown function/Ispoprotein, putative
SAG0961	247	+				+	-		sortase SrtA
SAG0963	320	+							conserved hypothetical protein
SAG0971	282	+		+		+	-		protein of unknown function/lipoprotein, putative
SAG0973	320	+						+	nısin-resistance protein, putative
SAG0977	312				+				sensor histidine kinase
SAG0979	553	+		+		+	•		ABC transporter, substrate-binding protein
SAG0984	437	+							sensor histidine kinase CiaH
SAG0992	286	+		+		+	+		phosphate ABC transporter, phosphate-binding protein
									iron-compound ABC transporter, iron-compound-binding
SAG1007	342	+		+		+	-		protein
SAG1014	190	+				-	-		conserved hypothetical protein
SAG1018	40			+				+	lipoprotein, putative
SAG1024	183	3 +		+					lipoprotein, putative
SAG1029	10	+							hypothetical protein
SAG1030	30	4 +				+	+		protein of unknown function
SAG1037	15	7 +						+	hypothetical protein
SAG1052	4	7	+	T				+	cell walf surface anchor family protein, putative
SAG1072	20	0 +							conserved hypothetical protein
SAG1094	27	8			+	+	+		conserved hypothetical protein
									spermidine/putrescine ABC transporter,
SAG1108	35	7 +				+	-		spermidine/putrescine-binding prot.
SAG1121	29	5 +			T		1	1	polysaccharide dencetylase family protein
SAG1120	22	8 +	1		1	+	+	1	protein of unknown function
SAG112	7 44	6 +				1		+	conserved domain protein
SAG1130	4	9 +	1				$\top$	+	hypothetical protein
SAG113	3 6	4 +		1	1				conserved hypothetical protein
SAG113	19	3 +	1		1	1	1	1	conserved hypothetical protein

	Size	Signal	Sortase	Lipo-		Western		GBS	
ORF	(aa)	Peptide	motif	protein	Other	blot	FACS	specific	Annotation
SAG1149	207	+		+					lipoprotein, putative
SAG1184	236	+							conserved hypothetical protein
SAG1186	553				+				metallo-beta-lactamase superfamily protein
SAG1189	334	+							conserved hypothetical protein
SAG1190	551				+				adherence and virulence protein A
SAG1197	1072	+							hyaluronidase
SAG1201	367	+							iminodiacetate oxidase, putative
SAG1206	854	+							conserved domain protein
SAG1214	58	+							hypothetical protein
SAG1216	1252		+			+	-		pullulanase, putative
SAG1227	198	+				+	-		protein of unknown function
SAG1233	822	+				+	-		streptococcal histidine triad family protein
SAG1234	306	+		+		+	+		lamınin-bindıng surface protem
SAG1238	202	+							hypothetical protein
SAG1283	1631		+			+	+		agglutinin receptor
SAG1313	56	+							conserved hypothetical protein
SAG1327	409	+							sensor histidine kinase
SAG1331	979	+	+			+	+		R5 protein
SAG1333	690	+	+			+	+		5'-nucleotidase family protein
SAG1350	544	+						,	surface antigen-related protein
SAG1361	414	+							conserved hypothetical protein
SAG1371	392	+							conserved hypothetical protein
SAG1393	310			+					iron compound ABC transporter, substrate-binding protein
SAG1404	30	3 +	+			+	-		cell wall surface anchor family protein
SAG1405	294	1 +			+	+	+		sortase family protein
SAG1406	29	3 +							sortase family protein
SAG1407	70:	5 +	+		1	+	+		cell wall surface anchor family protein
SAG1408	90	1	+						cell wall surface anchor family protein
SAG1419	57	7		+			T	+	lipoprotein, putative
SAG1431	26	8		+			$\top$		amino acid ABC transporter, amino acid-binding protein
SAG1433	37.	5 +	1	1					conserved hypothetical protein
				1					. maltose/maltodextrin ABC transporter,
SAG1441	41.	5 +				+	+		maltose/maltodextrin-binding protem



	Size	Signal	Sortase	Lipo-		Western		GBS	
ORF	(aa)	Peptide	motif	protein	Other	blot	FACS	specific	Annotation
SAG1462	970		+						cell wall surface anchor family protein
SAG1473	192	+	+					+	cell wall surface anchor family protein
SAG1474	680	+	+						amidase family protein
SAG1483	78	+							preprotein translocase, SecG subunit
SAG1488	195	+				+	+		dephospho-CoA kmase
SAG1491	530	+						+	hypothetical protein
SAG1508	590				+	+	•		67 kDa Myosin-crossreactive streptococcal antigen
SAG1518	538	+		+					peptide ABC transporter, peptide-binding protein
SAG1530	267	+		+		+	-		peptidyl-prolyl cis-trans isomerase, cyclophilin-type
									manganese ABC transporter, manganese-binding adhesion
SAG1533	308	+		+		+	•	ļ	Isprotein
SAG1544	232	+							gluconate 5-dehydrogenase, putative
SAG1551	67	+						+	hypothetical protein
SAG1552	719	+							conserved hypothetical protein
SAG1553	477	+						+	hypothetical protein
SAG1562	280	+							conserved hypothetical protein
									branched-chain ammo acid ABC transporter, ammo acid-
SAG1582	388	+		+		+	-		binding protem
SAG1590	449				+	+	+		potassium uptake protem, Trk family
SAG1601	79	+							conserved hypothetical protein
SAG1610	28	5		+		+	-		amino acid ABC transporter, substrate-binding protein
SAG1618	103	2			+	+	+		Snf2 family protein
SAG1624	50	1 +							sensor histidine kinase CsrS
SAG1628	18	4 +							lemA protein
SAG1631	22	3 +				+	1 -		potassium uptake protein, Trk family, putative
SAG1641	27	4 +				+	-		YaeC family protein
SAG1642	27	7 +	1	+		+	1	1	ABC transporter, substrate-binding protein
SAG1683	51	2 +	1						immunogenic secreted protem, putative
SAG1706	23	8 +				1			conserved hypothetical protein
SAG1745	14	8 +					1	+	hypothetical protein
SAG1752	39	0 +			1	1	1		conserved hypothetical protein TIGR00275
SAG1759	23	0		1	<del>                                     </del>	+	+	1	protein of unknown function
SAG1762	16	9 +	1	1	1	1	+		conserved hypothetical protein

	Size	Signal	Sortase	Lipo-		Western		GBS	
ORF	(aa)	Peptide	motif	protein	Other	blot	FACS	specific	Annotation
SAG1767	289	+		+					acid phosphatase
SAG1768	336			<u> </u>	+	+	+		glyceraldehyde 3-phosphate dehydrogenase
SAG1774	424	+			<del> </del>				conserved hypothetical protein
SAG1786	130	+				+	•		protein of unknown function
SAG1787	420	+							dltD protein
SAG1791	395	+							sensor histidine kinase
SAG1822	272	+				+	-		protein of unknown function
SAG1823	418				+	+	+		protein of unknown function
SAG1837	468				+				prophage LambdaSa2, lysin, putative
SAG1838	109	+		1					prophage LambdaSa2, holin, putative
SAG1839	136	+							conserved hypothetical protein
SAG1842	1224				+				prophage LambdaSa2, PblB, putative
SAG1912	194	+							N-acetylmuramoyl-L-alanine amidase, family 4 protein
SAG1921	508	3 +			,				sensor histidine kinase
SAG1932	810	5 +							neuraminidase-related protein
SAG1938	30	7 +		+		+	1		adhesion lipoprotein
SAG1941	80	0 +	+			+	-		2',3'-cyclto-nucleotide 2'-phosphodiesterase
SAG1945	34:	5 +							iron ABC transporter, iron-binding protein
SAG1947	54	9			+				conserved hypothetical protein
SAG1960	55	1			+	+	+		sensor histidine kinase
SAG1966	29	3		+		+	-		hemolysin precursor, putative
SAG1996	26	3 +	+				T		cell wall surface anchor family protein, putative
SAG1997	18	2 +							hypothetical protein
SAG1998	45	7 +							hypothetical protein
SAG202	82	6	+						cell wall surface anchor family protein
SAG2043	25	5 +			T				cAMP factor
SAG2053	3 157	70 +	+		1				scrine protease, subtilase family, putative
SAG205	5 46	52			+	T-			sensor histidine kinase
SAG205	5 20	)2 +						+	chromosome assembly-related protein
SAG206	3 63	30 +	+						pathogenicity protein, putative
SAG207	8 32	20 +		+	T	+	-		protein of unknown function/lipoprotein, putative
	T						1		competence/damage-inducible protein CmA, authentic
SAG209	4	+				+	+		frameshift

	Size	Signal	Sortase	Lipo-		Western		GBS	
ORF	(aa)	Peptide	motif	protein	Other	blot	FACS	specific	Annotation
SAG2121	223	+						+	hypothetical protein
SAG2123	454	+							sensor histidine kinase
SAG2141	660	+				+	-		DHH family protein
SAG2147	234	+		+		+	+		protein of unknown function/hipoprotein, putative
SAG2148	179	+	<b></b>						LysM domain protein
SAG2174	409	+						<b>-</b>	serine protease
SAG0013	428	+				+	-		protein of unknown function



ORF	Ann tation
SAG0038	conserved hypothetical protein
SAG0048	transcriptional regulator Cro/CI family
SAG0091	transcriptional regulator ComX1 putative
SAG0137	conserved hypothetical protein
SAG0686	DNA-entry nuclease putative
SAG0770	membrane protein putative
SAG0868	DNA-entry nuclease
SAG1143	conserved hypothetical protein
SAG1233	streptococcal histidine triad family protein
SAG1596	integrase/recombinase phage integrase family
SAG1616	conserved hypothetical protein
SAG1721	conserved hypothetical protein.

Probable recently duplicated genes are indicated on the same line and are separated by a semicolon.

SAG0148 oligopeptide ABC transporter, substrate-binding protein, putative; SAG0979 ABC transporter, substrate-binding protein

SAG0151 oligopeptide ABC transporter, ATP-binding protein; SAG1515 peptide ABC transporter, ATP-binding protein

SAG0195 IS1548, transposase; SAG0693 IS1548, transposase; SAG0760 IS1548, transposase; SAG0945 IS1548, transposase; SAG1584

IS1548, transposase; SAG1619 IS1548, transposase

SAG0230 conserved hypothetical protein; SAG1039 conserved hypothetical protein

SAG0233 hypothetical protein; SAG1785 hypothetical protein

SAG0261 IS1381, transposase OrfB; SAG0542 IS1381, transposase OrfA; SAG0543 IS1381, transposase OrfB; SAG0966 IS1381, transposase

OrfB; SAG1457 IS1381, transposase OrfB; SAG1550 IS1381, transposase OrfB; SAG2002 IS1381, transposase OrfB

SAG0262 IS1381, transposase OrfA; SAG0965 IS1381, transposase OrfA; SAG1549 IS1381, transposase OrfA; SAG1458 IS1381, transposase

OrfA; SAG2003 IS1381, transposase OrfA

SAG0383 protein of unknown function/lipoprotein, putative; SAG0785 conserved hypothetical protein

SAG0405 protein of unknown function/lipoprotein, putative; SAG0954 protein of unknown function/lipoprotein, putative

SAG0417 glycosyl transferase, group 2 family protein; SAG1422 glycosyl transferase, group 2 family protein

SAG0429 oxidoreductase, aldo/keto reductase family; SAG1476 oxidoreductase, aldo/keto reductase family

SAG0432 transcriptional regulator, AraC family; SAG0644 transcriptional regulator, AraC family

SAG0434 transposase, IS256 family, truncation; SAG0448 transposase, IS256 family

SAG0438 bacteriophage L54a, integrase, truncation; SAG1986 site-specific recombinase, phage integrase family; SAG1989 hypothetical

protein; SAG1993 site-specific recombinase, phage integrase family; SAG2115 hypothetical protein

SAG0442 acetyltransferase, GNAT family; SAG0443 acetyltransferase, GNAT family

SAG0447 magnesium transporter, CorA family; SAG0875 magnesium transporter, CorA family, putative

SAG0508 beta-lactam resistance factor; SAG1349 beta-lactam resistance factor

SAG0566 prophage LambdaSa1, single-strand binding protein; SAG1713 single-strand binding protein; SAG1863 prophage LambdaSa2, single-

strand binding protein

SAG0603 conserved hypothetical protein; SAG1838 prophage LambdaSa2, holin, putative

SAG0604 prophage LambdaSa1, lysin, putative; SAG1837 prophage LambdaSa2, lysin, putative

SAG0618 transposase OrfB, IS3 family, truncation; SAG0639 transposase OrfB, IS3 family; SAG1232 transposase OrfB, IS3 family,

truncation; SAG1242 transposase OrfB, IS3 family, truncation

SAG0640 transposase OrfA, IS3 family; SAG1241 transposase OrfA, IS3 family

SAG0646 cell wall surface anchor family protein; SAG1404 cell wall surface anchor family protein

SAG0647 sortase family protein; SAG0648 sortase family protein; SAG0650 sortase family protein

SAG0649 cell wall surface anchor family protein, putative; SAG1408 cell wall surface anchor family protein

SAG0676 proteinase, putative; SAG2053 serine protease, subtilase family, putative

SAG0679 protein of unknown function; SAG0680 protein of unknown function; SAG0681 conserved domain protein

SAG1002 protease, putative; SAG1465 protease, putative

SAG1025 hypothetical protein; SAG1033 FtsK/SpoIIIE family protein

SAG1067 IS861, transposase OrfA; SAG1526 IS861, transposase OrfA

SAG1068 IS861, transposase OrfB; SAG1256 IS861, transposase OrfB, truncation; SAG1527 IS861, transposase OrfB

SAG1140 conserved hypothetical protein; SAG1141 conserved hypothetical protein

SAG1164 glycosyl transferase CpsJ(V); SAG1165 glycosyl transferase CpsO(V)

SAG1182 phosphopentomutase; SAG2069 phosphopentomutase

SAG1225 conserved hypothetical protein; SAG1540 conserved hypothetical protein

SAG1228 ISSdy1, transposase OrfA; SAG1243 ISSdy1, transposase OrfA

SAG1229 ISSdy1, transposase OrfB; SAG1244 ISSdy1, transposase OrfB

SAG1253 transposase, ISL3 family; SAG2022 transposase, ISL3 family

SAG1254 mercuric reductase; SAG2023 mercuric reductase

SAG1255 mercuric resistance operon regulatory protein MerR; SAG2024 mercuric resistance operon regulatory protein MerR

SAG1259 conserved hypothetical protein; SAG1272 conserved hypothetical protein

SAG1283 agglutinin receptor; SAG2021 cell wall surface anchor family protein

SAG1297 C-5 cytosine-specific DNA methylase; SAG1869 prophage LambdaSa2, type II DNA modification methyltransferase, putative

SAG1405 sortase family protein; SAG1406 sortase family protein

SAG1414 glycosyl transferase, group 2 family protein; SAG1415 glycosyl transferase, group 2 family protein

SAG1456 glycosyl transferase, family 8, degenerate; SAG2060 glycosyl transferase, family 8

SAG1521 transposase, IS30 family, putative; SAG1576 transposase, IS30 family, putative, truncation; SAG1795 transposase, IS30 family,

putative

SAG1655 transcriptional regulator, MerR family; SAG1972 transcriptional regulator, MerR family

SAG1979 membrane protein, putative; SAG2034 membrane protein, putative

SAG1980 ABC transporter, ATP-binding protein; SAG2035 ABC transporter, ATP-binding protein

SAG1982 transcriptional regulator, Cro/CI family; SAG2037 transcriptional regulator, Cro/CI family

SAG1983 conserved hypothetical protein; SAG2039 conserved hypothetical protein

Table 4: Probable recently duplicated genes

SAG1984 conserved hypothetical protein TIGR00730; SAG2040 conserved hypothetical protein TIGR00730

SAG1988 conserved hypothetical protein; SAG2114 conserved hypothetical protein

Strain	Source	Capsular serotype	Reference
090	Lancefield	Ia	
515	Houston	Ia	(1)
A909	Lancefield	Ia	(2)
Davis	Channing	Ia	
DK1	Houston	Ia	
DK8	Houston	· Ia	
H36b	Lancefield	Ib	(2)
(S7) 7357b	Channing	Ib	(3)
18RS21	Lancefield	п	(4)
DK21	Houston	п	
СОН1	Seattle	Ш	(5)
СОН31	Seattle	Ш	(6)
D136C	Lancefield	III	(4)
M781	Houston	III	(7)
M732	Houston	III	(8)
1169NT1	Atlanta	v	(9)
2603V/R	Italy	v	This study
СЈВ111	Houston	v	(10)
JM9130013	Japan	VIII	(11)
SMU014	Japan	VIII	(11)
CJB110	Houston	Nontypeable	(12)



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Cluster 1	
SAG0230	conserved hypothetical protein
SAG0231	hypothetical protein
SAG0232	hypothetical protein
SAG0233	hypothetical protein
SAG0234	hypothetical protein
SAG0235	hypothetical protein
Cluster 2	
SAG0222	conserved domain protein
SAG0223	conserved hypothetical protein, fusion
SAG0225	hypothetical protein
SAG0226	recombination protein
SAG0227	hypothetical protein
SAG0228	conserved hypothetical protein
SAG0229	conserved hypothetical protein
Cluster 3	
SAG0634	hypothetical protein
SAG0635	acid phosphatase, class B
SAG0636	conserved hypothetical protein
SAG0638	cell wall surface anchor family protein, interruption-N
SAG0640	transposase OrfA IS3 family

SAG0642	hypothetical protein
SAG0643	chaperonin, 33 kDa, degenerate
SAG0644	transcriptional regulator, AraC family
SAG0645	cell wall surface anchor family protein
SAG0646	cell wall surface anchor family protein
SAG0647	sortase family protein
SAG0648	sortase family protein
SAG0649	cell wall surface anchor family protein, putative
SAG0650	sortase family protein
SAG0651	protein of unknown function

# Cluster 4

SAG1898	PTS system, IID component
SAG1899	PTS system, IIC component
SAG1900	PTS system, IIB component
SAG1901	glucuronyl hydrolase
SAG1902	PTS system, IIA component
SAG1905	conserved hypothetical protein
SAG1906	carbohydrate kinase, PfkB family

# Cluster 5

SAG0247 hypothetical protein
SAG0248 hypothetical protein

pothetical protein  pothetical protein  atative secreted protein  roteinase, putative
utative secreted protein
•
roteinase, putative
ypothetical protein
rotein of unknown function
onserved domain protein
ABC transporter, ATP-binding protein
onserved hypothetical protein
S1381, transposase OrfB
S1381, transposase OrfA
S1381, transposase OrfA
S1381, transposase OrfB
S1381, transposase OrfB
conserved hypothetical protein
hypothetical protein
hypothetical protein
protein of unknown function
Protont of mudio tanonous

SAG1032	conserved hypothetical protein
Cluster 8	
SAG1253	transposase, ISL3 family
SAG1254	mercuric reductase
SAG1255	mercuric resistance operon regulatory protein MerR
SAG2022	transposase, ISL3 family
SAG2023	mercuric reductase
SAG2024	mercuric resistance operon regulatory protein MerR
Cluster 9	•
SAG1993	site-specific recombinase, phage integrase family
SAG1994	conserved hypothetical protein
SAG1995	hypothetical protein
SAG1996	cell wall surface anchor family protein, putative
SAG1997	hypothetical protein
SAG1998	hypothetical protein
SAG2000	membrane protein, putative
SAG2001	conjugal transfer protein, interruption-C
SAG2007	conserved hypothetical protein
SAG2008	conserved hypothetical protein
SAG2009	conserved hypothetical protein
SAG2010	hypothetical protein



SAG2011	conserved hypothetical protein
SAG2012	hypothetical protein
SAG2016	hypothetical protein
SAG2017	transcriptional regulator, Cro/CI family
SAG2025	Mn2+/Fe2+ transporter, NRAMP family
Cluster 10	
SAG1039	conserved hypothetical protein
SAG1447	conserved hypothetical protein
SAG1448	glycosyl transferase, group 1 family protein
SAG1449	preprotein translocase SecA subunit, putative
SAG1450	conserved domain protein
SAG1452	conserved hypothetical protein
SAG1453	preprotein translocase SecY family protein
SAG1454	glycosyl transferase, putative
SAG1455	glycosyl transferase, group 2 family protein
SAG1456	glycosyl transferase, family 8, degenerate
SAG1459	glycosyl transferase family 8
SAG1460	glycosyl transferase, family 8
SAG1461	conserved hypothetical protein
SAG1462	cell wall surface anchor family protein
SAG1463	transcriptional regulator, RofA family, authentic point mutation
SAG1469	conserved hypothetical protein

SAG1471	conserved hypothetical protein .
SAG1933	PTS system, IIC component, putative
Cluster 11	
SAG0009	hypothetical protein
SAG0120	hypothetical protein
SAG0157	deoxyribonuclease-related protein, degenerate
SAG0186	hypothetical protein
SAG0216	hypothetical protein
SAG0236	hypothetical protein
SAG0307	hypothetical protein
SAG0308	ABC transporter, ATP-binding protein
SAG0311	DNA-binding response regulator, authentic point mutation
SAG0518	peptide chain release factor 2, programmed frameshift
SAG0553	hypothetical protein
SAG0555	prophage LambdaSa1, antirepressor, putative
SAG0564	conserved hypothetical protein
SAG0579	conserved hypothetical protein
SAG0580	conserved hypothetical protein, truncation
SAG0611	transposase, degenerate
SAG0637	transcriptional regulator, TetR family, putative, authentic frameshif
SAG0641	Tn5252, Orf 10 protein, degenerate
SAG0652	Tn5252, Orf 28 protein, degenerate

SAG0655	conserved hypothetical protein
SAG0678	endopeptidase O, degenerate
SAG0683	transmembrane protein Vexp3, putative, degenerate
SAG0855	glycogen biosynthesis protein GlgD, authentic frameshift
SAG0898	hypothetical protein
SAG0899	hypothetical protein
SAG0901	hypothetical protein
SAG0902	hypothetical protein
SAG0903	hypothetical protein
SAG0917	Tn916, hypothetical protein
SAG0920	Tn916, hypothetical protein
SAG0922	Tn916, hypothetical protein
SAG0924	Tn916, tetM leader peptide
SAG0928	Tn916, hypothetical protein, authentic frameshift
SAG0936	Tn916, hypothetical protein
SAG0943	hypothetical protein
SAG0972	conserved hypothetical protein, authentic frameshift
SAG1023	hypothetical protein
SAG1080	hypothetical protein
SAG1123	hypothetical protein
SAG1129	hypothetical protein
SAG1136	conserved hypothetical protein
SAG1217	conserved hypothetical protein, authentic frameshift



SAG1231	transposase OrfB, IS3 family, degenerate
SAG1242	transposase OrfB, IS3 family, truncation
SAG1309	hypothetical protein
SAG1331	R5 protein
SAG1437	hypothetical protein
SAG1445	MutT/nudix family protein, authentic frameshift
SAG1484	ribosomal protein L33
SAG1493	hypothetical protein
SAG1539	hypothetical protein
SAG1543	conserved hypothetical protein, authentic frameshift
SAG1560	hypothetical protein
SAG1568	phosphoserine aminotransferase, authentic frameshift
SAG1570	conserved hypothetical protein
SÁG1601	conserved hypothetical protein
SAG1644	hypothetical protein
SAG1646	hypothetical protein
SAG1699 .	hypothetical protein
SAG1705	peptidase, M24 family, authentic point mutation
SAG1708	hypothetical protein
SAG1857	prophage LambdaSa2, HNH endonuclease family protein
SAG1864	hypothetical protein
SAG1868	hypothetical protein

SAG1869	prophage LambdaSa2, type II DNA modification methyltransferase,	
putative	·	
SAG1872	hypothetical protein .	
SAG1874	hypothetical protein	
SAG1876	prophage LambdaSa2, HNH endonuclease family protein	
SAG1878	conserved domain protein	
SAG1881	hypothetical protein	
SAG1883	conserved hypothetical protein	
SAG1886	hypothetical protein	
SAG1903	hypothetical protein	
SAG1937	streptococcal histidine triad family protein, degenerate	
SAG1971	hypothetical protein	
SAG1979	membrane protein, putative	
SAG1980	ABC transporter, ATP-binding protein	
SAG1981	hypothetical protein	
SAG1982	transcriptional regulator, Cro/CI family	
SAG1983	conserved hypothetical protein	
SAG1984	conserved hypothetical protein TIGR00730	
SAG1985	hypothetical protein	
SAG1991	transcriptional regulator, Cro/CI family	
SAG1992	protein of unknown function	
SAG1999	hypothetical protein	
SAG2004	conjugal transfer protein, interruption-N	

SAG2039	conserved hypothetical protein
SAG2044	hypothetical protein
SAG2052	hypothetical protein
SAG2065	ribosomal protein L33
SAG2094	competence/damage-inducible protein CinA, authentic frameshift
SAG2099	hypothetical protein
Cluster 12	
SAG1164	glycosyl transferase CpsJ(V)
SAG1165	glycosyl transferase CpsO(V)
SAG1166	glycosyl transferase CpsN(V)
SAG1167	polysaccharide biosynthesis protein CpsM(V)
SAG1168	polysaccharide biosynthesis protein cpsH(V)
Cluster 13	
SAG0581	conserved hypothetical protein
SAG0582	conserved hypothetical protein
· SAG0583	conserved hypothetical protein
SAG0585	conserved hypothetical protein
SAG0586	conserved hypothetical protein
SAG0587	prophage LambdaSal, structural protein, putative
SAG0588	conserved hypothetical protein
SAG0589	conserved hypothetical protein

SAG0590	conserved hypothetical protein
SAG0591	conserved hypothetical protein
SAG0593	prophage LambdaSa1, structural protein
SAG0594	conserved hypothetical protein
SAG0595	conserved hypothetical protein
SAG0596	prophage LambdaSa1, pblA protein, internal deletion
Cluster 14	
SAG0915	Tn916, transposase
SAG0918	Tn916, hypothetical protein
SAG0919	Tn916, hypothetical protein
SAG0921	Tn916, transcriptional regulator, putative
SAG0925	Tn916, hypothetical protein
SAG0926	Tn916, NLP/P60 family protein
SAG0927	membrane protein, putative
SAG0929	Tn916, hypothetical protein
SAG0930	Tn916, hypothetical protein
SAG0931	Tn916, hypothetical protein
SAG0932	Tn916, transcriptional regulator, putative
SAG0933	Tn916, FtsK/SpoIIIE family protein
SAG0934	Tn916, hypothetical protein
SAG0935	Tn916, hypothetical protein
SAG0937	ABC transporter, ATP-binding protein, authentic frameshift

Cluster 15	
SAG1835	conserved hypothetical protein
SAG1837	prophage LambdaSa2, lysin, putative
SAG1839	conserved hypothetical protein
SAG1840	hypothetical protein
SAG1842	prophage LambdaSa2, PblB, putative
SAG1843	conserved hypothetical protein
SAG1844	conserved hypothetical protein
SAG1849	hypothetical protein
SAG1851	conserved domain protein
SAG1852	conserved domain protein
SAG1853	prophage LambdaSa2, protease, putative
SAG1854	conserved hypothetical protein
SAG1855	prophage LambdaSa2, terminase large subunit, putative
SAG1856	hypothetical protein
SAG1858	hypothetical protein
SAG1859	prophage LambdaSa2, site-specific recombinase, phage integrase family
SAG1860	conserved hypothetical protein
SAG1861	prophage LambdaSa2, transcriptional regulator, Cro/CI family
SAG1862	hypothetical protein
SAG1863	prophage LambdaSa2, single-strand binding protein
SAG1865	conserved hypothetical protein

SAG1866	conserved hypothetical protein
SAG1867	conserved hypothetical protein
SAG1870	prophage LambdaSa2, DNA replication protein DnaC, putative
SAG1871	prophage LambdaSa2, bacteriophage replication protein/hypothetical
protein, trunc	eation/fusion
SAG1873	prophage LambdaSa2, replicative DNA helicase
SAG1877	prophage LambdaSa2, antirepressor protein, putative
SAG1879	hypothetical protein
SAG1882	prophage LambdaSa2, repressor protein, putative
SAG1884	hypothetical protein
SAG1885	prophage LambdaSa2, site-specific recombinase, phage integrase family
Cluster 16	•
SAG1247	site-specific recombinase, phage integrase family
SAG1250	Tn5252, relaxase
SAG1251	Tn5252, Orf 9 protein
SAG1252	Tn5252, Orf 10 protein
SAG1256	IS861, transposase OrfB, truncation
SAG1257	cation-transporting ATPase, E1-E2 family
SAG1258	cadmium efflux system accessory protein
SAG1259	conserved hypothetical protein
SAG1260	hypothetical protein
SAG1261	conserved hypothetical protein

SAG1262	cation-transporting ATPase, E1-E2 family
SAG1263	conserved domain protein, authentic frameshift
SAG1264	transcriptional repressor CopY, putative
SAG1265	cadmium resistance transporter, putative
SAG1266	hypothetical protein
SAG1267	hypothetical protein
SAG1268	repressor protein, putative
SAG1270	ImpB/MucB/SamB family protein
SAG1271	conserved hypothetical protein
SAG1272	conserved hypothetical protein
SAG1273	conserved hypothetical protein
SAG1274	conserved hypothetical protein
SAG1276	conserved hypothetical protein
SAG1277	hypothetical protein
SAG1278	hypothetical protein
SAG1279	conserved domain protein
SAG1280	SNF2 family protein
SAG1281	hypothetical protein
SAG1283	agglutinin receptor
SAG1284	abortive infection protein AbiGI .
SAG1285	abortive infection protein AbiGII
SAG1286	Tn5252, Orf28
SAG1287	Tn5252, Orf26

SAG1288	Tn5252, Orf25, degenerate
SAG1289	Tn5252, Orf23
SAG1290	hypothetical protein
SAG1291	Tn5252, Orf 21 protein, internal deletion
SAG1292	hypothetical protein
SAG1293	protease, putative
SAG1294	conserved hypothetical protein
SAG1295	conserved hypothetical protein
SAG1296	conserved hypothetical protein
SAG1297	C-5 cytosine-specific DNA methylase
SAG1299	conserved hypothetical protein
SAG1304	hypothetical protein

Locus Annotation

Housekeeping

SAG0466 thiolase

SAG0471 glucokinase

SAG0492 amino acid ABC transporter, ATP-binding protein

SAG0767 D-alanine--D-alanine ligase

SAG1086 xanthine phosphoribosyltransferase

SAG1600 glutamate racemase

SAG1680 shikimate 5-dehydrogenase

SAG1723 · signal peptidase I

### Surface-exposed

SAG0079 adenylate kinase

SAG0093 D-alanyl-D-alanine carboxypeptidase family protein

SAG0163 competence protein CglA

SAG0290 ABC transporter, substrate-binding protein

SAG0368 protein of unknown function

SAG0503 lipase/acylhydrolase

SAG1473 cell wall surface anchor family protein

SAG1552 conserved hypothetical protein

SAG1641 YaeC family protein

SAG2147 protein of unknown function/lipoprotein, putative

SAG2148 LysM domain protein

ORFxxxxx Annotation
ORF00003 PcsB protein (pscB)
ORF00004 ribose-phosphate pyrophosphokinase (prsA)
ORF00005 aminotransferase, class I
ORF00006 recombination protein O
ORF00009 fatty acid/phospholipid synthesis protein PlsX (plsX)
ORF00011 phosphoribosylaminoimidazole-succinocarboxamide synthase (purC)
ORF00012 phosphoribosylformylglycinamidine synthase, putative
ORF00013 amidophosphoribosyltransferase (purF)
ORF00014 phosphoribosylformylglycinamidine cyclo-ligase (purM)
ORF00015 phosphoribosylglycinamide formyltransferase (purN)
ORF00020 group B streptococcal surface immunogenic protein
ORF00021 N-acetylmannosamine-6-P epimerase, putative
ORF00022 sugar ABC transporter, sugar-binding protein
ORF00023 sugar ABC transporter, permease protein
ORF00023 sugar ABC transporter, permease protein
ORF00024 sugar ABC transporter, permease protein
ORF00027 N-acetylneuraminate lyase, putative
ORF00027 N-acetyfiletramilate lyase, putative
ORF00028 expressed ROK lamily protein ORF00030 phosphosugar-binding transcriptional regulator, RpiR family, putative
ORF00031 phosphoribosylamine—glycine ligase (purD) ORF00032 phosphoribosylaminoimidazole carboxylase, catalytic subunit (purE)
ORF00032 phosphoribosylaminoimidazole carboxylase, Catalytic suburit (purE)  ORF00033 phosphoribosylaminoimidazole carboxylase, ATPase subunit (purK)
ORF00036 adenylosuccinate lyase (purB)
ORF00037 transcriptional regulator, Cro/Cl family
ORF00038 Holliday junction DNA helicase RuvB (ruvB)
ORF00039 phosphotyrosine protein phosphatase, low molecular weight
ORF00040 MORN motif family protein
ORF00041 membrane protein, putative
ORF00043 alcohol dehydrogenase, propanol-preferring (adhP)
ORF00045 MATE efflux family protein
ORF00046 ribosomal protein S10 (rpsJ)
ORF00047 ribosomal protein L3 (rplC)
ORF00048 ribosomal protein L4 (rpID)
ORF00049 ribosomal protein L23 (rpIW)
ORF00050 ribosomal protein L2 (rplB)
ORF00052 ribosomal protein S19 (rpsS)
ORF00054 ribosomal protein L22 (rplV)
ORF00055 ribosomal protein S3 (rpsC)
ORF00056 ribosomal protein L16 (rplP)
ORF00058 ribosomal protein L29 (rpmC)
ORF00059 ribosomal protein S17 (rpsQ)
ORF00060 ribosomal protein L14 (rplN)
ORF00061 ribosomal protein L24 (rplX)
ORF00063 ribosomal protein L5 (rplE)
ORF00065 ribosomal protein S8 (rpsH)
ORF00066 ribosomal protein L6 (rplF)
ORF00068 ribosomal protein L18 (rpIR)
ORF00069 ribosomal protein S5 (rpsE)
ORF00070 ribosomal protein L30 (rpmD)
ORF00071 ribosomal protein L15 (rplO)
ORF00072 preprotein translocase, SecY subunit
ORF00073 adenylat kinase (adk)
ORF00074 translation initiation factor IF-1 (infA)
ORF00075 ribosomal protein L36 (rpmJ)
ORF00077 ribosomal protein S13 (rpsM)
[2 22 Harden, B. 2.0 (Pa)

ORFxxxxx Annotati n
ORF00078 ribosomal protein S11 (rpsK)
ORF00080 DNA-directed RNA polymerase, alpha subunit (rpoA)
ORF00093 transcriptional regulator ComX1, putative
ORF00094 phosphoglycerate mutase family protein
ORF00097 heat-inducible transcription repressor HrcA (hrcA)
ORF00098 heat shock protein GrpE (grpE)
ORF00099 dnaK protein (dnaK)
ORF00100 dnaJ protein (dnaJ)
ORF00101 transcriptional regulator, GntR family
ORF00102 tRNA pseudouridine synthase A (truA)
ORF00103 phosphomethylpyrimidine kinase, putative
ORF00104 conserved hypothetical protein
ORF00105 conserved hypothetical protein
ORF00106 conserved hypothetical protein
ORF00107 trigger factor (tig)
ORF00108 DNA-directed RNA polymerase, delta subunit, putative
ORF00109 CTP synthase (pyrG)
ORF00111 deoxyuridine 5'-triphosphate nucleotidohydrolase (dut)
ORF00113 carbonic anhydrase-related protein
ORF00115 pyridine nucleotide-disulphide oxidoreductase family protein
ORF00116 glutamyl-tRNA synthetase (gltX)
ORF00119 ribose ABC transporter, ATP-binding protein (rbsA)
ORF00122 ribose operon repressor RbsR (rbsR)
ORF00125 ABC transporter, ATP-binding protein
ORF00126 DNA-binding response regulator
ORF00128 sensor histidine kinase
ORF00131 fructose-bisphosphate aldolase (fba)
ORF00132 L-2-hydroxyisocaproate dehydrogenase
ORF00133 ribosomal protein L28 (rpmB)
ORF00134 conserved hypothetical protein
ORF00135 DAK2 domain protein
ORF00136 expressed SPFH domain/Band 7 family protein
ORF00141 amino acid ABC transporter, ATP-binding protein
ORF00142 amino acid ABC transporter, amino acid-binding protein/permease protein
ORF00143 conserved hypothetical protein
ORF00145 undecaprenol kinase, putative
ORF00146 negative regulator of competence MecA, putative
ORF00149 ABC transporter, ATP-binding protein
ORF00150 conserved hypothetical protein
ORF00151 selenocysteine lyase (csdB)
ORF00152 NifU family protein
ORF00153 conserved hypothetical protein
ORF00155 D-alanyl-D-alanine carboxypeptidase
ORF00158 oligopeptide ABC transporter, permease protein
ORF00160 oligopeptide ABC transporter, ATP-binding protein
ORF00161 oligopeptide ABC transporter, ATP-binding protein
ORF00167 adc operon repressor AdcR (adcR)
ORF00168 zinc ABC transporter, ATP-binding protein
ORF00169 zinc ABC transporter, permease protein
ORF00172 tyrosyl-tRNA synth tase (tyrS)
ORF00173 penicillin-binding protein 1B, putative
ORF00174 DNA-directed RNA polymerase, beta subunit (rpoB)
ORF00176 DNA-directed RNA polymerase beta' subunit (rpoC)
ORF00178 conserved hypothetical protein
ORF00179 competenc protein CgIA (cgIA)
•

ORFxxxxx Ann tati n
ORF00180 competence protein CglB (cglB)
ORF00181 conserved hypothetical protein
ORF00183 conserved hypothetical protein
ORF00184 acetate kinase (ackA)
ORF00190 pyrroline-5-carboxylate reductase (proC)
ORF00191 glutamyl-aminopeptidase (pepA)
ORF00198 single-strand binding protein (ssb)
ORF00211 PTS system, IIABC components
ORF00212 alpha amylase family protein
ORF00214 transcriptional antiterminator, BgIG family
ORF00219 PTS system, IIC component, putative
ORF00224 ribosomal protein S15 (rpsO)
ORF00225 polyribonucleotide nucleotidyltransferase (pnp)
ORF00227 serine O-acetyltransferase (cysE)
ORF00229 cysteinyl-tRNA synthetase (cysS)
ORF00230 conserved hypothetical protein
ORF00231 RNA methyltransferase, TrmH family, group 3
ORF00233 DegV family protein
ORF00236 ribosomal protein L13 (rplM)
ORF00237 ribosomal protein S9 (rpsl)
ORF00261 transcriptional regulator MutR family
ORF00262 transporter, putative
ORF00263 amino acid ABC transporter, permease protein
ORF00264 amino acid ABC transporter, amino acid-binding protein
ORF00265 amino acid ABC transporter, permease protein
ORF00266 amino acid ABC transporter, ATP-binding protein
ORF00295 N-acetylglucosamine-6-phosphate deacetylase (nagA)
ORF00296 conserved hypothetical protein
ORF00297 glycyl-tRNA synthetase, alpha subunit (glyQ)
ORF00299 glycyl-tRNA synthetase, beta subunit (glyS)
ORF00300 conserved hypothetical protein
ORF00302 glycerol kinase (glpK)
ORF00303 alpha-glycerophosphate oxidase
ORF00304 glycerol uptake facilitator protein (glpF)
ORF00306 conserved hypothetical protein
ORF00307 transketolase (tkt)
ORF00309 ABC transporter, ATP-binding protein
ORF00310 membrane protein, putative
ORF00313 PTS system, IIBC components
ORF00314 glutamate 5-kinase (proB)
ORF00315 gamma-glutamyl phosphate reductase (proA)
ORF00316 conserved hypothetical protein TIGR00006
ORF00318 penicillin-binding protein 2X (pbpX)
ORF00319 phospho-N-acetylmuramoyl-pentapeptide-transferase (mraY)
ORF00319 phospho-N-acetylindramoyi-pentapeptide-transferase (mra r) ORF00320 ATP-dependent RNA helicase, DEAD/DEAH box family
ORF00321 ABC transporter, substrate-binding protein
ORF00321 ABC transporter, substrate-binding protein  ORF00322 amino acid ABC transporter, permease protein
ORF00323 amino acid ABC transporter, ATP-binding protein
ORF00325 thloredoxin reductase (trxB)
ORF00326 conserved hypothetical protein ORF00327 NAD synthetas (nadE)
ORF00328 aminopeptidase C (pepC) ORF00329 penicillin-binding prot in 1A (pbp1A)
ORF00330 recombination protein U (recU)
ORF00331 conserved hypothetical prot_in
ON 0000 I Conserved hypothetical prot in

ORFxxxxx Annotation
ORF00335 conserved hypothetical protein
ORF00336 conserved hypothetical protein
ORF00337 autoinducer-2 production protein LuxS (luxS)
ORF00338 KH domain protein
ORF00348 guanylate kinase (gmk)
ORF00349 DNA-directed RNA polymerase, omega subunit, putative
ORF00350 primosomal protein N' (priA)
ORF00351 methionyl-tRNA formyltransferase (fmt)
ORF00352 Sun protein (sun)
ORF00353 serine/threonine phosphatase, putative
ORF00354 serine/threonine protein kinase
ORF00355 conserved hypothetical protein
ORF00356 sensor histidine kinase, putative
ORF00358 DNA-binding response regulator
ORF00359 hydrolase, haloacid dehalogenase family/peptidyl-prolyl cis-trans isomerase, cyclophilin type
ORF00360 general stress protein, putative
ORF00361 pyruvate formate-lyase-activating enzyme (pflA)
ORF00362 transcriptional regulator, DeoR family
ORF00363 transcriptional regulator, putative
ORF00364 PTS system, cellobiose-specific IIA component (celC)
ORF00366 PTS system, cellobiose-specific IIB component (celA)
ORF00367 PTS system, cellobiose-specific IIC component (celB)
ORF00368 formate acetyltransferase (pflD)
ORF00369 transaldolase family protein
ORF00371 glycerol dehydrogenase (gldA)
ORF00372 cysteine synthase A (cysK)
ORF00373 conserved hypothetical protein TIGR00257
ORF00374 helicase, putative
ORF00374 helicase, putative ORF00375 competence protein F, putative
ORF00376 ribosomal subunit interface protein (yfiA)
ORF00376 hibosoffial subulific frace protein (yil/x) ORF00385 enoyl-CoA hydratase/isomerase family protein
ORF00386 transcriptional regulator, MarR family
ORF00387 3-oxoacyl-(acyl-carrier-protein) synthase III (fabH)
ORF00388 acyl carrier protein (acpP)
ORF00390 enoyl-(acyl-carrier-protein) reductase II (fabK)
ORF00390 enbyl-(acyl-carrier-protein) reductase it (labit) ORF00391 malonyl CoA-acyl carrier protein transacylase (fabD)
ORF00392 3-oxoacyl-[acyl-carrier protein] reductase (fabG)
ORF00393 3-oxoacyl-(acyl-carrier-protein) synthase II (fabF)
ORF00394 acetyl-CoA carboxylase, biotin carboxyl carrier protein (accB)
ORF00395 (3R)-hydroxymyristoyl-(acyl-carrier-protein) dehydratase (fabZ)
ORF00396 acetyl-CoA carboxylase, biotin carboxylase (accC)
ORF00397 acetyl-CoA carboxylase, carboxyl transferase, beta subunit (accD)
ORF00398 acetyl-CoA carboxylase, carboxyl transferase, alpha subunit (accA)
ORF00400 seryl-tRNA synthetase (serS)
ORF00403 conserved hypothetical protein
ORF00404 PTS system, mannose-specific IID component
ORF00405 PTS system, mannose-specific IIC component (manM)
ORF00406 PTS system, mannose-specific IIAB components (manL)
ORF00407 hydrolase, haloacid dehalogenase-like family
ORF00410 xanthine/uracil permease family protein
ORF00411 conserved hypothetical protein TIGR00150, putative
ORF00412 acetyltransferas , GNAT family
ORF00413 expressed protein of unknown function
ORF00415 HIT family protein (hit)
ORF00419 ABC transporter, ATP-binding protein

ORFxxxxx Annotation
ORF00421 ABC transporter, permease protein
ORF00422 conserved hypothetical protein
ORF00423 conserved hypothetical protein TIGR00091
ORF00424 conserved hypothetical protein, POINT MUTATION
ORF00425 N utilization substance protein A (nusA)
ORF00426 conserved hypothetical protein
ORF00427 ribosomal protein L7A family
ORF00428 translation initiation factor IF-2
ORF00429 ribosome-binding factor A (rbfA)
ORF00432 copper-transporter ATPase CopA
ORF00435 hydrolase, haloacid dehalogenase-like family
ORF00436 DNA polymerase I (polA)
ORF00437 CoA binding domain protein
ORF00440 DNA-binding response regulator
ORF00441 sensor histidine kinase
ORF00443 queuine tRNA-ribosyltransferase (tgt)
ORF00444 conserved hypothetical protein
ORF00449 glucose-6-phosphate isomerase (pgi)
ORF00451 rhomboid family protein
ORF00457 Mornibold family protein
ORF00453 UTP-glucose-1-phosphate uridylyltransferase (galU)
ORF00454 glycerol-3-phosphate dehydrogenase (NAD(P)+) (gpsA)
ORF00455 ribonuclease P protein component (mpA)
ORF00456 SpollIJ family protein
ORF00458 R3H domain protein
ORF00463 conserved hypothetical protein
ORF00464 RecX protein
ORF00465 RNA methyltransferase, TrmA family
ORF00470 ribonucleoside-diphosphate reductase 2, beta subunit (nrdF)
ORF00472 ribonucleoside-diphosphate reductase 2, alpha subunit (nrdE)
ORF00482 alcohol dehydrogenase, zinc-containing
ORF00483 oxidoreductase, aldo/keto reductase family
ORF00484 cation efflux system protein
ORF00485 transcriptional regulator, TetR family
ORF00496 conserved hypothetical protein
ORF00500 acetyltransferase, GNAT family
ORF00501 conserved hypothetical protein
ORF00502 valyI-tRNA synthetase (valS)
ORF00508 aspartate—ammonia ligase (asnA)
ORF00511 type II DNA modification methyltransferase, putative
ORF00513 phosphopantetheine adenylyltransferase (coaD)
ORF00515 conserved hypothetical protein
ORF00519 conserved hypothetical protein
ORF00520 conserved hypothetical protein TIGR00048
ORF00522 ABC transporter, ATP-binding/permease protein
ORF00522 ABC transporter, ATP-binding/permease protein
ORF00524 anthranilate synthase component II (trpG)
ORF00532 endonuclease III (nth)
ORF00532 endonuclease iii (ntn) ORF00534 conserved hypothetical protein
ORF00535 glucokinase (glk)
ORF00536 expressed prot in with rhodan se domain
ORF00537 elongation factor Tu family protein
ORF00540 UDP-N-acetylmuramoylalanineD-glutamate ligase (murD)
ORF00541 UDP-N-acetylglucosamine-N-acetylmuramyl-(pentapeptide) pyrophosphoryl-undecaprenol N-
acetylglucosamine transferase (murG)

ORFxxxxx Annotati n
ORF00542 cell division protein DivIB, putative
ORF00544 cell division protein FtsA (ftsA)
ORF00545 cell division protein FtsZ (ftsZ)
ORF00546 ylmE protein, putative
ORF00547 ylmF protein (ylmF)
ORF00549 ylmH protein (ylmH)
ORF00550 cell division protein DivIVA, putative
ORF00552 isoleucyl-tRNA synthetase (ileS)
ORF00553 conserved hypothetical protein
ORF00554 MutT/nudix family protein
ORF00555 ATP-dependent Clp protease, ATP-binding subunit
ORF00557 conserved hypothetical protein
ORF00558 amino acid ABC transporter, permease protein
ORF00559 amino acid ABC transporter, ATP-binding protein
ORF00560 phosphoglucomutase/phosphomannomutase family protein
ORF00562 methylenetetrahydrofolate dehydrogenase/methenyltetrahydrofolate cyclohydrolase (folD)
ORF00564 exodeoxyribonuclease VII, large subunit (xseA)
ORF00566 geranyltranstransferase, putative
ORF00567 hemolysin A
ORF00570 DNA repair protein RecN (recN)
ORF00571 expressed DegV family protein
ORF00574 DNA-binding protein HU (hup)
ORF00576 dihydroorotate dehydrogenase A (pyrDA)
ORF00577 beta-lactam resistance factor (fibB)
ORF00578 beta-lactam resistance factor (fibA)
ORF00579 murM protein, putative
ORF00580 hydrolase, haloacid dehalogenase-like family
ORF00581 HD domain protein
ORF00582 conserved hypothetical protein
ORF00583 cation-transporting ATPase, E1-E2 family
ORF00588 cell division ABC transporter, ATP-binding protein FtsE (ftsE)
ORF00589 cell division ABC transporter, permease protein FtsX (ftsX)
ORF00591 metallo-beta-lactamase superfamily protein
ORF00593 DNA polymerase III, epsilon subunit/ATP-dependent helicase DinG
ORF00595 aspartate aminotransferase (aspC)
ORF00596 asparaginyl-tRNA synthetase (asnS)
ORF00601 conserved hypothetical protein
ORF00602 conserved hypothetical protein
ORF00603 conserved hypothetical protein
ORF00605 zinc ABC transporter, zinc-binding adhesion liprotein
ORF00606 ribosomal protein L31 (rpmE)
ORF00607 DHH family protein
ORF00609 flavodoxin
ORF00614 ribosomal protein L19 (rplS)
ORF00640 prophage LambdaSa1, single-strand binding protein (ssb)
ORF00693 DNA-binding response regulator VncR (vncR)
ORF00694 sensor histidine kinase VncS (vncS)
ORF00699 rod shape-determining protein RodA, putative□ (rodA)
ORF00700 hydrolase, haloacid dehalogenase-like family
ORF00701 DNA gyrase, B subunit (gyrB)
ORF00702 septation ring formation regulator EzrA, putative
ORF00705 conserved hypothetical protein
ORF00706 enolase (eno)
ORF00708 3-phosphoshikimate 1-carboxyvinyltransferase (aroA)
ORF00709 shikimate kinas (aroK)

ORF00710 psr protein	
ORF00711 RNA methyltransferase, T	rmA family
ORF00729 sortase family protein	
ORF00731 sortase family protein	
ORF00734 sortase family protein, FR	AMESHIFT
ORF00743 ABC transporter, ATP-bind	ding protein
ORF00744 membrane protein	
ORF00745 conserved hypothetical pro	otein
ORF00748 cylG protein (cylG)	
ORF00776 DNA-entry nuclease, puta	tive
ORF00789 2-keto-3-deoxygluconate I	kinase
ORF00792 2-dehydro-3-deoxyphosph	nogluconate aldolase/4-hydroxy-2-oxoglutarate aldolase (eda)
ORF00798 proline dipeptidase (pepQ	
ORF00799 transcriptional regulator, F	RegM family
ORF00802 glycosyl transferase, grou	p 1 family protein
ORF00803 threonyl-tRNA synthetase	(thrS)
ORF00804 DNA-binding response reg	gulator
ORF00808 amino acid ABC transport	er, permease protein
ORF00810 amino acid ABC transport	er, ATP-binding protein
ORF00811 DNA-binding response re-	gulator
ORF00812 sensory box histidine kina	se
ORF00813 metallo-beta-lactamase fa	mily protein
ORF00815 ribonuclease III (rnc)	
ORF00816 expressed putative chrom	
ORF00817 hydrolase, haloacid dehal	
ORF00818 hydrolase, haloacid dehal	
ORF00819 signal recognition particle	
ORF00820 ABC transporter, substrat	
ORF00821 ABC transporter, permeas	
ORF00824 transcriptional accessory	
ORF00825 conserved hypothetical pr	
ORF00828 HPr(Ser) kinase/phospha	
ORF00830 prolipoprotein diacylglyce	
ORF00832 conserved hypothetical properties of the conserved hypothetical prope	
ORF00835 peptidase, U32 family, pu	tative
ORF00836 peptidase, U32 family	
ORF00837 conserved hypothetical p	
ORF00844 lysyl-tRNA synthetase (ly	sS) ·
ORF00846 phosphoglycerate mutase	
ORF00847 ebsC family protein, puta	tive
ORF00850 peptidase, U32 family	
ORF00855 oligoendopeptidase F, pu	
ORF00856 phosphoenolpyruvate car	
ORF00859 cell division protein, FtsW	//RodA/SpoVE family (ftsW)
ORF00861 translation elongation fac	
ORF00863 triosephosphate isomera:	se (tpiA)
ODEOGOOF -basels :	

ORF00869 UDP-N-acetylmuramoylalanyl-D-glutamyl-2,6-diaminopimelate--D-alanyl-D-alanyl ligase (murF)

ORF00865 phosphoglycerate mutase (gpmA)
ORF00867 recombination protein RecR (recR)
ORF00868 D-alanine—D-alanine ligase

ORF00873 peptide chain releas factor 3 (prfC)
ORF00876 ABC transporter, ATP-binding protein

ORF00880 ATP-dependent RNA helicase, DEAD/DEAH box family

ORF00870 oxalate:formate antiporter ORF00871 membrane protein, putative

ORFxxxxx Annotati n
ORF00882 conserved hypothetical protein
ORF00883 conserved hypothetical protein
ORF00884 acyltransferase family protein
ORF00885 competence protein CelA (celA)
ORF00887 DNA internalization-related competence protein ComEC/Rec2
ORF00889 sugar-binding transcriptional regulator, Lacl family
ORF00892 DNA polymerase III, delta subunit, putative
ORF00893 superoxide dismutase, Fe-Mn (sodA)
ORF00894 transcriptional antiterminator LicT
ORF00895 PTS system, beta-glucosides-specific IIABC components
ORF00896 6-phospho-beta-glucosidase (bglA)
ORF00899 glycerate kinase 2 (garK)
ORF00904 S-adenosylmethionine:tRNA ribosyltransferase-isomerase (queA)
ORF00906 glucosamine-6-phosphate isomerase (nagB)
ORF00908 ribosomal small subunit pseudouridine synthase
ORF00911 competence protein CoiA (coiA)
ORF00911 competence protein ColA (ColA)  ORF00912 oligoendopeptidase B (pepB)
ORF00914 O-methyltransferase family protein
ORF00916 protease maturation protein, putative
ONFOUS 15 didniyrthing synthetase (alab)
ORF00925 transcriptional regulator, Cro/CI family
ORF00928 ribonucleoside-diphosphate reductase 2, beta subunit (nrdF)
ORF00929 ribonucleoside-diphosphate reductase 2, alpha subunit (nrdE)
ORF00930 ribonucleoside-diphosphate reductase 2, NrdH-redoxin (nrdH)
ORF00931 phosphocarrier protein HPr (ptsH)
ORF00932 phosphoenolpyruvate-protein phosphotransferase (ptsl)
ORF00933 glyceraldehyde-3-phosphate dehydrogenase, NADP-dependent (gapN)
ORF00934 polysaccharide deacetylase family protein
ORF00935 ATP-dependent RNA helicase, DEAD/DEAH box family
ORF00936 uridine kinase (udk)
ORF00937 conserved hypothetical protein
ORF00938 DNA polymerase III, gamma and tau subunits (dnaX)
ORF00940 biotin—acetyl-CoA-carboxylase ligase
ORF00941 S-adenosylmethionine synthetase (metK)
ORF00955 UDP-N-acetylglucosamine 1-carboxyvinyltransferase (murA)
ORF00956 acetyltransferase, GNAT family
ORF00957 CBS domain protein
ORF00958 methionine aminopeptidase, type I (map)
ORF00959 ribonuclease BN, putative
ORF00962 conserved hypothetical protein
ORF00963 DNA ligase, NAD-dependent (ligA)
ORF00964 BmrU protein, putative
ORF00966 pullulanase, putative
ORF00973 ATP synthase F0, A subunit (atpB)
ORF00974 ATP synthase F0, B subunit (atpF)
ORF00975 ATP synthase F1, delta subunit (atpH)
ORF00976 ATP synthase F1, alpha subunit (atpA)
ORF00977 ATP synthase F1, gamma subunit (atpG)
ORF00978 ATP synthase F1, beta subunit (atpD)
ORF00979 ATP synthase F1, epsilon subunit (atpC)
ORF00981 UDP-N-acetylglucosamin 1-carboxyvinyltransferase (murA)
ORF00983 DNA-entry nuclease (endA)
ORF00984 phenylalanyl-tRNA synthetas , alpha subunit (pheS)
ORF00986 phenylalanyl-tRNA synthetas , beta subunit (pheT)
ORF00988 exonuclease RexB (rexB)
OTHER DESIGNATIONS (1971)

ORFxxxxx Ann tati n
ORF00989 exonucleas RexA (rexA)
ORF00991 tRNA modification GTPase TrmE (trmE)
ORF00992 ABC transporter, ATP-binding protein
ORF00993 acetoin dehydrogenase, thymine PPi dependent, E1 component, alpha subunit
ORF00994 acetoin dehydrogenase, thymine PPi dependent, E1 component, beta subunit
ORF00995 acetoin dehydrogenase, thymine PPi dependent, E2 component, dihydrolipoamide
ORF00996 acetoin dehydrogenase, thymine PPi dependent, E3 component, dihydrolipoamide dehydrogenase
ORF00997 lipoate-protein ligase A (IpIA)
ORF00998 cobyric acid synthase, putative
ORF00999 mur ligase family protein
ORF01000 conserved hypothetical protein TIGR00159
ORF01001 expressed protein of unknown function
ORF01002 phosphoglucomutase/phosphomannomutase family protein
ORF01005 oxygen-independent coproporphyrinogen III oxidase, putative
ORF01006 conserved hypothetical protein
ORF01007 hydrolase, haloacid dehalogenase-like family
ORF01008 conserved hypothetical protein
ORF01023 GTP-binding protein LepA (lepA)
ORF01027 PilB-related protein
ORF01030 cation-transporting ATPase, E1-E2 family
ORF01033 conserved hypothetical protein
ORF01040 Tn916, tetracycline resistance protein (tetM)
ORF01057 transcriptional regulator, GntR family
ORF01058 DNA polymerase III, alpha subunit (dnaE)
ORF01059 6-phosphofructokinase (pfk)
ORF01060 pyruvate kinase (pyk)
ORF01063 glucosaminefructose-6-phosphate aminotransferase (isomerizing) (glmS)
ORF01066 phnA protein (phnA)
ORF01068 amino acid ABC transporter, permease protein
ORF01069 amino acid ABC transporter, ATP-binding protein
ORF01070 amino acid ABC transporter, amino acid-binding protein
ORF01072 ribosomal protein S20 (rpsT)
ORF01073 pantothenate kinase (coaA)
ORF01074 conserved hypothetical protein
ORF01075 cytidine deaminase (cdd)
ORF01076 expressed putative lipoprotein
ORF01077 sugar ABC transporter, ATP-binding protein
ORF01078 sugar ABC transporter, permease protein, putative
ORF01079 sugar ABC transporter, permease protein, putative
ORF01080 NADH oxidase (nox-2)
ORF01081 L-lactate dehydrogenase (ldh)
ORF01082 DNA gyrase, A subunit (gyrA)
ORF01083 sortase SrtA (srtA)
ORF01089 GMP synthase (guaA)
ORF01090 transcriptional regulator, GntR family
ORF01091 gld protein (gid)
ORF01093 expressed putative lipoprotein
ORF01097 ABC transporter, ATP-binding protein
ORF01099 DNA-binding response regulator
ORF01101 site-specific recombinase, phage integrase family
ORF01106 signal recognition particle protein Ffh (ffh)
ORF01108 conserved hypothetical protein
ORF01109 sensor histidine kinase CiaH
ORF01110 DNA-binding response regulator CiaR (ciaR)
ORF01111 aminopeptidas N (pepN)
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ORFxxxxx Annotation
ORF01112 phosphate transport system regulatory protein PhoU (phoU)
ORF01113 phosphate ABC transporter, ATP-binding protein PstB, putative
ORF01114 phosphate ABC transporter, ATP-binding protein PstB, putative
ORF01115 phosphate ABC transporter, permease protein PstA, putative
ORF01116 phosphate ABC transporter, permease protein
ORF01117 phosphate ABC transporter, phosphate-binding protein
ORF01118 NOL1/NOP2/sun family protein
ORF01119 inositol monophosphatase family protein
ORF01120 conserved hypothetical protein
ORF01121 conserved hypothetical protein
ORF01122 macrolide-efflux protein mreA/riboflavin biosynthesis protein RibF
ORF01123 tRNA pseudouridine synthase B (truB)
ORF01125 conserved hypothetical protein
ORF01128 permease, putative
ORF01129 ABC transporter, ATP-binding protein
ORF01131 DNA topoisomerase I (topA)
ORF01132 DprA/SMF protein, putative DNA processing factor (dprA)
ORF01134 iron compound ABC transporter, ATP-binding protein
ORF01137 acetyltransferase, CysE/LacA/LpxA/NodL family
ORF01138 ribonuclease HII (rnhB)
ORF01139 GTP-binding protein
ORF01176 carbamoyl-phosphate synthase, large subunit (carB)
ORF01177 carbamoyl-phosphate synthase, small subunit (carA)
ORF01178 aspartate carbamoyltransferase (pyrB)
ORF01179 dihydroorotase, multifunctional complex type (pyrC)
ORF01180 orotate phosphoribosyltransferase (pyrE)
ORF01181 orotidine 5'-phosphate decarboxylase (pyrF)
ORF01183 ABC transporter, ATP-binding protein
ORF01184 ribonucleotide reductase, truncation
ORF01188 cardiolipin synthetase (cls)
ORF01189 formatetetrahydrofolate ligase (fhs)
ORF01190 lipoate-protein ligase A (IpIA)
ORF01198 flavoprotein-related protein
ORF01199 flavoprotein family protein
ORF01200 membrane protein, putative
ORF01201 phosphoglucomutase (pgm)
ORF01203 IS861, transposase OrfB
ORF01205 ABC transporter, ATP-binding/permease protein
ORF01206 ABC transporter, ATP-binding/permease protein
ORF01207 conserved hypothetical protein
ORF01208 conserved hypothetical protein
ORF01209 Serine hydroxymethyltransferase
ORF01210 Sua5/YciO/YrdC/YwlC family protein
ORF01211 modification methylase, HemK family
ORF01212 peptide chain release factor 1 (prfA)
ORF01213 thymidine kinases (tdk)
ORF01214 4-oxalocrotonate tautomerase (xyIM)
ORF01216 ApbE family protein
ORF01220 xanthine permease (pbuX)
ORF01221 xanthine phosphoribosyltransferase (xpt)
ORF01222 guanosine monophosphate reductase (guaC)
ORF01227 phosphate acetyltransferase
ORF01227 phosphate acetyltransierase ORF01228 ribosomal large subunit pseudouridine synthase, RluD subfamily
ORF01229 express d protein of unknown function
ORF01230 GTP pyrophosphokinase family protein
OTA OTZOO OTT Pyrophosphokindse latility protein

### ORFxxxxx Annotati n ORF01231 conserved hypothetical protein ORF01232 ribose-phosphate pyrophosphokinase (prsA) ORF01233 cysteine desulphurase (iscS) ORF01234 conserved hypothetical protein ORF01235 conserved hypothetical protein ORF01236 DNA repair protein RadC (radC) ORF01238 6-phospho-beta-glucosidase (ascB) ORF01239 platelet activating factor, putative ORF01240 hydrolase, haloacid dehalogenase-like family ORF01242 voltage-gated chloride channel family protein ORF01243 spermidine/putrescine ABC transporter, spermidine/putrescine-binding protein (potD) ORF01244 spermidine/putrescine ABC transporter, permease protein (potC) ORF01245 spermidine/putrescine ABC transporter, permease protein (potB) ORF01246 spermidine/putrescine ABC transporter, ATP-binding protein (potA) ORF01247 UDP-N-acetylenolpyruvoylglucosamine reductase (murB) ORF01248 2-amino-4-hydroxy-6-hydroxymethyldihydropteridine pyrophosphokinase (folk) ORF01250 dihydropteroate synthase (folP) ORF01251 GTP cyclohydrolase I (folE) ORF01252 folylpolyglutamate synthase (folC) ORF01259 aldehyde dehydrogenase family protein ORF01260 membrane protein ORF01274 gls24 protein, putative ORF01276 gls24 protein, putative ORF01279 conserved hypothetical protein ORF01282 ATP-dependent DNA helicase PcrA (pcrA) ORF01283 conserved hypothetical protein, FRAMESHIFT ORF01284 uracil permease (uraA) ORF01285 sodium:alanine symporter family protein ORF01286 cation efflux family protein ORF01290 ribosomal protein S1 (rpsA) ORF01292 branched-chain amino acid aminotransferase (ilvE) ORF01294 DNA topoisomerase IV, A subunit (parC) ORF01295 DNA topoisomerase IV, B subunit (parE) ORF01296 membrane protein, putative ORF01297 uracil-DNA glycosylase (ung) ORF01317 transcriptional regulator, LysR family, putative ORF01319 purine nucleoside phosphorylase (deoD) ORF01321 purine nucleoside phosphorylase (deoD) ORF01323 phosphopentomutase (deoB) ORF01324 ribose 5-phosphate isomerase (rpiA) ORF01327 tributyrin esterase (estA) ORF01328 metallo-beta-lactamase superfamily protein ORF01329 ABC transporter, ATP-binding protein ORF01330 ABC transporter, permease protein ORF01331 conserved hypothetical protein ORF01332 adherence and virulence protein A (pavA) ORF01335 TPR domain protein ORF01336 membrane protein ORF01338 mutator MutT protein (mutX) ORF01339 hyaluronidase ORF01343 iminodiacetate oxidas, putative ORF01344 conserved hypothetical protein TIGR00486 ORF01345 conserved hypothetical protein ORF01346 DNA replication protein Dnad, putativ

ORF01347 adenine phosphoribosyltransferase (apt)

ORFxxxxx Annotation
ORF01350 single-stranded-DNA-specific xonuclease RecJ (recJ)
ORF01351 oxidoreductase, short chain dehydrogenase/reductase family
ORF01352 metallo-beta-lactamase superfamily protein
ORF01353 conserved hypothetical protein
ORF01354 GTP-binding protein HflX (hflX)
ORF01355 tRNA delta(2)-isopentenylpyrophosphate transferase (miaA)
ORF01357 exfoliative toxin A, putative
ORF01358 pullulanase, putative
ORF01362 conserved hypothetical protein
ORF01363 peptidase, M20/M25/M40 family
ORF01364 nitroreductase family protein
ORF01367 excinuclease ABC, C subunit (uvrC)
ORF01380 streptococcal histidine triad family protein
ORF01381 laminin-binding surface protein (lmb)
ORF01397 Tn5252, relaxase
ORF01403 mercuric reductase (merA)
ORF01406 IS861, transposase OrfB, truncation
ORF01407 cation-transporting ATPase, E1-E2 family
ORF01411 conserved hypothetical protein
ORF01412 cation-transporting ATPase, E1-E2 family
ORF01415 transcriptional repressor CopY, putative
ORF01416 cadmium resistance transporter, putative
ORF01451 C-5 cytosine-specific DNA methylase
ORF01453 conserved hypothetical protein
ORF01455 ribosomal protein L7/L12 (rplL)
ORF01456 ribosomal protein L10 (rplJ)
ORF01458 ATP-dependent Clp protease, ATP-binding subunit
ORF01467 GTP-binding protein (cgpA)
ORF01468 ATP-dependent Clp protease, ATP-binding subunit ClpX (clpX)
ORF01470 dihydrofolate reductase (folA)
ORF01471 thymidylate synthase (thyA)
ORF01472 HMG-CoA synthase
ORF01473 3-hydroxy-3-methylglutaryl-CoA reductase
ORF01474 conserved hypothetical protein
ORF01475 hemolysin III, putative
ORF01476 conserved hypothetical protein TIGR00147
ORF01479 isopentenyl-diphosphate delta-isomerase
ORF01480 phosphomevalonate kinase
ORF01481 diphosphomevalonate decarboxylase (mvaD)
ORF01482 mevalonate kinase, putative
ORF01484 DNA-binding response regulator
ORF01491 polypeptide deformylase, putative
ORF01495 ABC transporter, ATP-binding/permease protein
ORF01496 ABC transporter, ATP-binding/permease protein
ORF01498 ABC transporter, ATP-binding protein
ORF01499 polyA polymerase family protein
ORF01500 DegV family protein
ORF01501 expressed protein of unknown function
ORF01504 PTS system, fructose specific IIABC components
ORF01505 1-phosphofructokinase (fruK)
ORF01506 lactose phosphotransferase system repressor (lacR)
ORF01507 beta-lactam resistance factor
ORF01511 pyridine nucleotide-disulphide oxidoreductase family protein
ORF01512 tRNA (guanine-N1)-methyltransferase (trmD)
ORF01513 16S rRNA processing protein RimM (rimM)

ORFxxxxx Annotation
ORF01515 transcriptional regulator, RofA family
ORF01516 KH domain protein
ORF01517 ribosomal protein S16 (rpsP)
ORF01518 permease, putative
ORF01519 ABC transporter, ATP-binding protein
ORF01520 conserved hypothetical protein
ORF01523 carbamoyl-phosphate synthase, small subunit (carA)
ORF01524 pyrimidine operon regulatory protein (pyrR)
ORF01525 ribosomal large subunit pseudouridine synthase, RluD subfamily
ORF01526 lipoprotein signal peptidase (IspA)
ORF01527 transcriptional regulator, LysR family
ORF01528 ribosomal protein L27 (rpmA)
ORF01529 conserved hypothetical protein
ORF01530 ribosomal protein L21 (rplU)
ORF01531 conserved hypothetical protein, FRAMESHIFT
ORF01532 thiamine biosynthesis protein Thil (thil)
ORF01533 cysteine desulphurase (iscS)
ORF01536 glutathione reductase (gor)
ORF01537 conserved hypothetical protein
ORF01538 chorismate synthase (aroC)
ORF01539 3-dehydroquinate synthase (aroB)
ORF01540 3-dehydroquinate dehydratase (aroD)
ORF01541 conserved hypothetical protein
ORF01543 ribosomal protein L20 (rplT)
ORF01544 ribosomal protein L35 (rpml)
ORF01545 translation initiation factor IF-3 (infC)
ORF01546 cytidylate kinase (cmk)
ORF01548 ferredoxin, 4Fe-4S
ORF01550 peptidase t (pepT)
ORF01551 polysaccharide biosynthesis protein, putative
ORF01552 UDP-N-acetylmuramoylalanyl-D-glutamate2,6-diaminopimelate ligase (murE)
ORF01553 iron compound ABC transporter, ATP-binding protein (fepC)
ORF01555 iron compound ABC transporter, permease protein
ORF01556 iron compound ABC transporter, permease protein
ORF01558 inorganic pyrophosphatase, manganese-dependent (ppa)
ORF01559 pyruvate formate-lyase-activating enzyme (pflA) ORF01560 CBS domain protein
ORF01561 conserved hypothetical protein
ORF01564 PAP2 family protein
ORF01565 membrane protein, putative
ORF01567 expressed sortage family protein
ORF01568 sortase family protein
ORF01571 rogB protein FRAMESHIFT (rogB)
ORF01587 conserved hypothetical protein
ORF01589 RNA polymerase sigma-70 factor (rpoD) ORF01590 DNA primase (dnaG)
ORF01591 large conductance mechanosensitive channel protein (mscL)
ORF01592 ribosomal protein S21 (rpsU)
ORF01594 amino acid ABC transporter, amino acid-binding protein ORF01598 rhodanese family protein
ORF01602 glycogen phosphorylase (glgP)
ORF01603 4-alpha-glucanotransferase (malQ)
ORF01604 maltose operon repressor MalR, putative
ORF01605 maltose/maltodextrin ABC transporter, maltose/maltodextrin-binding protein
ORF01606 maltose ABC transporter, permease protein

ORFxxxxx Ann tati n
ORF01607 maltose ABC transporter, permease protein
ORF01614 preprotein translocase SecA subunit, putative
ORF01619 preprotein translocase SecY family protein
ORF01634 excinuclease ABC, B subunit (uvrB)
ORF01636 glutamine ABC transporter, glutamine-binding protein/permease protein (glnP)
ORF01637 glutamine ABC transporter, ATP-binding protein, GlnQ putative
ORF01640 GTP-binding protein, GTP1/Obg family (obg)
ORF01646 amidase family protein
ORF01647 ribosomal small subunit pseudouridine synthase A (rsuA)
ORF01648 oxidoreductase, aldo/keto reductase family
ORF01651 lactoylglutathione lyase (gloA)
ORF01652 glycosyl transferase, group 2 family protein
ORF01654 SsrA-binding protein (smpB)
ORF01655 exoribonuclease, VacB/Rnb family (vacB)
ORF01657 preprotein translocase, SecG subunit
ORF01658 multi-drug resistance protein
ORF01662 dephospho-CoA kinase
ORF01663 formamidopyrimidine-DNA glycosylase (mutM)
ORF01677 GTP-binding protein Era (era)
ORF01678 diacylglycerol kinase (dgkA)
ORF01679 conserved hypothetical protein TIGR00043
ORF01685 PhoH family protein
ORF01687 conserved hypothetical protein
ORF01689 conserved hypothetical protein
ORF01690 ribosome recycling factor (frr)
ORF01691 uridylate kinase (pyrH)
ORF01693 peptide ABC transporter, ATP-binding protein FRAMESHIFT
ORF01697 ribosomal protein L1 (rplA)
ORF01698 ribosomal protein L11 (rplK)
ORF01706 IS861, transposase OrfB
ORF01707 chorismate binding enzyme
ORF01708 FtsK/SpolIIE family protein
ORF01709 peptidyl-prolyl cis-trans isomerase, cyclophilin-type
ORF01710 manganese ABC transporter, permease protein
ORF01711 manganese ABC transporter, ATP-binding protein
ORF01712 manganese ABC transporter, manganese-binding adhesion liprotein
ORF01713 iron-dependent transcriptional regulator
ORF01714 5-methylthioadenosine nucleosidase/S-adenosylhomocysteine nucleosidase (pfs)
ORF01716 MutT/nudix family protein
ORF01718 UDP-N-acetylglucosamine pyrophosphorylase (glmU)
ORF01722 oxidoreductase, Gfo/ldh/MocA family
ORF01725 gluconate 5-dehydrogenase, putative
ORF01726 conserved hypothetical protein
ORF01738 branched-chain amino acid transport system II carrier protein (brnQ)
ORF01739 methionyl-tRNA synthetase (metG)
ORF01745 exodeoxyribonuclease (exoA)
ORF01746 conserved hypothetical protein
ORF01752 copper homeostasis protein CutC, putative
ORF01755 tetrapyrrole methylase family protein
ORF01756 conserved hypothetical protein
ORF01758 DNA polymerase III, delta prime subunit, putative
ORF01759 thymidylate kinase (tmk)
ORF01773 ATP-dependent Clp protease, proteolytic subunit ClpP (clpP)
ORF01774 uracil phosphoribosyltransferase (upp)
ORF01777 RNA methyltransferase, TrmH family, group 2

ORFxxxxx Ann tati n
ORF01781 conserved hypothetical protein TIGR00278
ORF01782 ribosomal large subunit pseudouridine synthase B (rluB)
ORF01783 conserved hypothetical protein TIGR00281
ORF01784 conserved hypothetical protein
ORF01785 integrase/recombinase, phage integrase family
ORF01786 CBS domain protein
ORF01787 conserved hypothetical protein
ORF01788 HAM1 protein
ORF01789 glutamate racemase (murl)
ORF01791 membrane protein, putative
ORF01792 transcriptional regulator, biotin repressor family
ORF01793 membrane protein, putative
ORF01795 RNA methyltransferase, TrmH family
ORF01796 acylphosphatase
ORF01797 lipoprotein, putative
ORF01799 amino acid ABC transporter, permease protein
ORF01801 amidase family protein
ORF01802 transcription elongation factor GreA (greA)
ORF01803 conserved hypothetical protein
ORF01804 acetyltransferase, GNAT family
ORF01805 UDP-N-acetylmuramatealanine ligase (murC)
ORF01806 conserved hypothetical protein
ORF01808 expressed putative helicase
ORF01811 phosphoglycerate dehydrogenase-related protein
ORF01812 primosomal protein Dnal (dnal)
ORF01813 conserved hypothetical protein
ORF01814 conserved hypothetical protein TIGR00244
ORF01815 sensor histidine kinase CsrS (csrS)
· ORF01816 DNA-binding response regulator CsrR (csrR)
ORF01817 conserved hypothetical protein
ORF01818 heat shock protein HtpX (htpX)
ORF01820 lemA protein (lemA)
ORF01821 glucose-inhibited division protein B (gidB)
ORF01822 sodium transport family protein
ORF01823 potassium uptake protein, Trk family, putative
ORF01825 ABC transporter, ATP-binding protein
ORF01828 branched-chain amino acid transport system II carrier protein (bmQ)
ORF01829 alcohol dehydrogenase, zinc-containing (adh)
ORF01830 ABC transporter, permease protein
ORF01831 ABC transporter, ATP-binding protein
ORF01833 expressed YaeC family protein
ORF01834 ABC transporter, substrate-binding protein
ORF01835 glutamine amidotransferase, class I
ORF01837 conserved hypothetical protein TIGR01033
ORF01846 glycerol uptake facilitator protein (glpF)
ORF01849 conserved hypothetical protein
ORF01851 conserved hypothetical protein
ORF01852 iojap-related protein
ORF01854 conserved hypothetical protein TIGR00488
ORF01855 conserved hypothetical protein TIGR00482
ORF01856 conserved hypothetical protein TIGR00253
ORF01857 GTP-binding protein
ORF01858 hydrolase, haloacid dehalogenase-like family
ORF01860 glutamyl-tRNA(Gln) amidotransferase, B subunit (gatB)
ORF01861 glutamyl-tRNA(GIn) amidotransferase, A subunit (gatA)
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ORFxxxxx Annotati n
ORF01862 glutamyl-tRNA(Gln) amidotransferase, C subunit (gatC)
ORF01867 isochorismatase family protein
ORF01869 transcriptional regulator CodY, putative
ORF01870 aminotransferase, class I
ORF01871 universal stress protein family FRAMESHIFT
ORF01872 hydrolase, haloacid dehalogenase-like family
ORF01873 asparaginase family protein
ORF01874 shikimate 5-dehydrogenase (aroE)
ORF01876 ATP-dependent DNA helicase RecG (recG)
ORF01878 alanine racemase (alr)
ORF01879 holo-(acyl-carrier-protein) synthase (acpS)
ORF01881 preprotein translocase, SecA subunit (secA)
ORF01882 mannose-6-phosphate isomerase, class I (manA)
ORF01883 fructokinase (scrK)
ORF01885 PTS system IIABC components
ORF01886 sucrose-6-phosphate hydrolase (scrB)
ORF01887 sucrose operon repressor ScrR (scrR)
ORF01888 N utilization substance protein B (nusB)
ORF01889 conserved hypothetical protein
ORF01890 translation elongation factor P (efp)
ORF01900 cytidine/deoxycytidylate deaminase family protein
ORF01906 excinuclease ABC, A subunit (uvrA)
ORF01907 conserved hypothetical protein
ORF01908 magnesium transporter, CorA family (corA)
ORF01909 ribosomal protein S18 (rpsR)
ORF01910 single-strand binding protein (ssb)
ORF01911 ribosomal protein S6 (rpsF)
ORF01912 A/G-specific adenine glycosylase (mutY)
ORF01914 thioredoxin (trx)
ORF01915 PAP2 family protein
ORF01916 MutS2 family protein
ORF01917 conserved hypothetical protein
ORF01918 conserved hypothetical protein
ORF01919 ribonuclease HIII (rnhC)
ORF01920 signal peptidase I
ORF01921 helicase, putative
ORF01923 DNA-damage inducible protein P (dinP)
ORF01924 formate acetyltransferase (pflD)
ORF01926 conserved hypothetical protein
ORF01927 proteinase, putative, degenerate, FRAMESHIFT
ORF01929 glycerol uptake facilitator protein, putative
ORF01930 universal stress protein family
ORF01933 X-pro dipeptidyl-peptidase (pepX)
ORF01937 ABC transporter, ATP-binding protein CydC (cydC)
ORF01938 ABC transporter, ATP-binding protein CydD
ORF01945 conserved hypothetical protein TIGR00103
ORF01948 exonuclease
ORF01949 conserved hypothetical protein
ORF01950 conserved hypothetical protein TIGR00275
ORF01952 ribosomal protein S14 (rpsN)
ORF01957 O-sialoglycoprotein endopeptidase family protein
ORF01958 ribosomal-protein-alanine acetyltransferase, putative
ORF01960 expressed protein of unknown function
ORF01961 conserved hypothetical protein
ORF01962 metallo-beta-lactamase superfamily protein

ORFxxxxx Ann tati n
ORF01963 conserved hypothetical protein
ORF01964 glutamine synthetase, type I (glnA)
ORF01965 transcriptional regulator GlnR (glnR)
ORF01967 conserved hypothetical protein
ORF01969 phosphoglycerate kinase (pgk)
ORF01971 glyceraldehyde 3-phosphate dehydrogenase (gap)
ORF01972 translation elongation factor G (fusA)
ORF01973 ribosomal protein S7 (rpsG)
ORF01974 ribosomal protein S12 (rpsL)
ORF01975 pur operon repressor (purR)
ORF01976 HD domain protein
ORF01977 conserved hypothetical protein
ORF01978 conserved hypothetical protein
ORF01979 ribulose-phosphate 3-epimerase (rpe)
ORF01980 conserved hypothetical protein TIGR00157
ORF01983 dimethyladenosine transferase (ksgA)
ORF01985 primase-related protein
ORF01987 deoxyribonuclease, TatD family
ORF01992 dltD protein (dltD)
ORF01993 D-alanyl carrier protein (dltC)
ORF01994 dltB protein (dltB)
ORF01996 D-alanine-activating enzyme (dltA)
ORF01997 sensor histidine kinase
ORF01998 DNA-binding response regulator
ORF01999 ribosomal protein L34 (rpmH)
ORF02004 amino acid ABC transporter, ATP-binding protein
ORF02007 conserved hypothetical protein
ORF02008 transcriptional antiterminator, BglG family
ORF02017 sugar binding transcriptional regulator, Lacl family
ORF02018 transaldolase family protein
ORF02019 carbohydrate isomerase, AraD/FucA family
ORF02020 hexulose-6-phosphate isomerase, putative
ORF02021 hexulose-6-phosphate synthase, putative
ORF02022 PTS system, IIA component
ORF02023 PTS system, IIB component
ORF02024 transport protein SgaT, putative
ORF02027 adenylosuccinate synthetase (purA)
ORF02033 chaperonin, 33 kDa (hslO)
ORF02034 NifR3/Smm1 family protein
ORF02037 ATP-dependent Clp protease, ATP-binding subunit
ORF02038 transcriptional regulator CtsR (ctsR)
ORF02040 translation elongation factor Ts (tsf)
ORF02041 ribosomal protein S2 (rpsB)
ORF02043 alkyl hydroperoxide reductase, subunit F (ahpF) ORF02076 prophage LambdaSa2, single-strand binding protein (ssb)
ORFO2076 propriage LambdaSa2, single-strand binding protein (SSO)
ORF02082 prophage LambdaSa2, type II DNA modification methyltransferase, putative
ORF02086 prophage LambdaSa2, replicative DNA helicase (dnaC)
ORF02104 endopeptidase O (pepO)
ORF02110 polypeptide deformylase (def)
ORF02111 sugar binding transcriptional regulator RegR (regR)
ORF02112 conserved hypothetical protein
ORF02113 PTS system, IID compon лt
ORF02114 PTS system, IIC component
ORF02115 PTS system, IIB component
ORF02116 glucuronyl hydrolase

### **ORFxxxxx** Annotati n ORF02118 PTS system, IIA component ORF02120 oxidoreductase, short-chain dehydrogenase/reductase family ORF02121 conserved hypothetical protein ORF02122 carbohydrate kinase, PfkB family ORF02123 2-dehydro-3-deoxyphosphogluconate aldolase/4-hydroxy-2-oxoglutarate aldolase (eda) ORF02127 DNA polymerase III, alpha subunit, Gram-positive type ORF02129 prolyl-tRNA synthetase (proS) ORF02130 membrane-associated zinc metalloprotease, putative ORF02131 phosphatidate cytidylyltransferase (cdsA) ORF02132 undecaprenyl diphosphate synthase (uppS) ORF02133 preprotein translocase, YajC subunit (yajC) ORF02140 glucan 1,6-alpha-glucosidase (dexB) ORF02141 sugar ABC transporter, ATP-binding protein (msmK) ORF02142 helix-turn-helix domain protein, fis-type ORF02144 tagatose 1,6-diphosphate aldolase (lacD) ORF02145 tagatose-6-phosphate kinase (lacC) ORF02146 galactose-6-phosphate isomerase, LacB subunit (lacB) ORF02147 galactose-6-phosphate isomerase, LacA subunit (lacA) ORF02149 PTS system, IIC component, putative ORF02150 PTS system, IIB component, putative ORF02152 PTS system, IIA component, putative ORF02153 lactose phosphotransferase system repressor (lacR) ORF02157 adhesion lipoprotein ORF02158 expressed protein of unknown function TIGR00256 ORF02159 GTP pyrophosphokinase (relA) ORF02161 nrdl protein (nrdl) ORF02164 iron ABC transporter, iron-binding protein ORF02165 DNA-binding response regulator ORF02167 PTS system, IID component ORF02168 PTS system, IIC component ORF02174 ABC transporter, ATP-binding protein ORF02176 response regulator ORF02177 conserved hypothetical protein ORF02178 PTS system, IIABC components ORF02179 sensor histidine kinase ORF02180 phosphate regulon response regulator PhoB (phoB) ORF02182 phosphate ABC transporter, ATP-binding protein (pstB) ORF02183 phosphate ABC transporter, permease protein ORF02184 phosphate ABC transporter, permease protein ORF02188 conserved hypothetical protein TIGR00046 ORF02189 ribosomal protein L11 methyltransferase (prmA) ORF02197 conserved hypothetical protein ORF02199 ATPase, AAA family ORF02249 mercuric reductase (merA) ORF02272 DNA topology modulation protein FlaR, putative ORF02273 glycerol dehydrogenase, putative ORF02281 DNA-binding response regulator ORF02285 leucyl-tRNA synthetase (leuS) ORF02290 transcription antitermination protein NusG (nusG) ORF02293 penicillin-binding protein 2A (pbp2A) ORF02294 ribosomal large subunit pseudouridine synthase, RluD subfamily ORF02296 phosphopentomutase (deoB) ORF02297 deoxyribose-phosphate aldolase (deoC) ORF02300 uridine phosphorylase (udp) ORF02302 60 kda chaperonin (groEL)

ORFxxxxx Ann tation
ORF02303 chaperonin, 10 kDa (groES)
ORF02305 ABC transporter, ATP-binding protein
ORF02306 ABC transporter, permease protein
ORF02307 expressed putative lipoprotein
ORF02309 glyoxalase family protein
ORF02310 conserved hypothetical protein
ORF02311 anaerobic ribonucleoside-triphosphate reductase activating protein (nrdG)
ORF02312 acetyltransferase, GNAT family
ORF02315 anaerobic ribonucleoside-triphosphate reductase (nrdD)
ORF02318 conserved hypothetical protein
ORF02320 conserved hypothetical protein
ORF02321 conserved hypothetical protein
ORF02322 recA protein (recA)
ORF02325 DNA-3-methyladenine glycosylase I (tag)
ORF02327 Holliday junction DNA helicase RuvA (ruvA)
ORF02329 DNA mismatch repair protein HexB (hexB)
ORF02333 arginine repressor ArgR, putative
ORF02334 arginyl-tRNA synthetase (argS)
ORF02337 conserved hypothetical protein
ORF02338 conserved hypothetical protein
ORF02339 aspartyl-tRNA synthetase (aspS)
ORF02340 histidyl-tRNA synthetase (hisS)
ORF02342 ribosomal protein L33 (rpmG)
ORF02357 DNA-binding response regulator
ORF02359 membrane protein, putative
ORF02360 carbamate kinase (arcC)
ORF02361 ornithine carbamoyltransferase (argF)
ORF02364 amino acid ABC transporter, ATP-binding protein
ORF02365 amino acid ABC transporter, permease and amino acid-binding protein
ORF02370 membrane protein, putative
ORF02371 transcriptional regulator, TetR family, putative
ORF02373 ribosomal protein S4 (rpsD)
ORF02374 conserved hypothetical protein
ORF02375 replicative DNA helicase (dnaC)
ORF02376 ribosomal protein L9 (rpll)
ORF02377 DHH family protein
ORF02378 glucose inhibited division protein A (gidA)
ORF02380 tRNA (5-methylaminomethyl-2-thiouridylate)-methyltransferase (trmU)
ORF02381 L-serine dehydratase, iron-sulfur-dependent, beta subunit (sdhB)
ORF02382 L-serine dehydratase, iron-sulfur-dependent, alpha subunit (sdhA)
ORF02385 cobalt transport family protein
ORF02386 ABC transporter, ATP-binding protein
ORF02387 ABC transporter, ATP-binding protein, FRAMESHIFT
ORF02388 CDP-diacylglycerolglycerol-3-phosphate 3-phosphatidyltransferase (pgsA)
ORF02389 peptidase, M16 family
ORF02390 conserved hypothetical protein
ORF02391 conserved hypothetical protein
ORF02392 recF protein (recF)
ORF02396 inosine-5'-monophosphate dehydrogenase (guaB)
ORF02397 transcriptional regulator, ArgR family
ORF02400 arginine deiminase (arcA)
ORF02402 ornithine carbamoyltransferase (argF)
ORF02404 carbamate kinas (arcC)
ORF02405 tryptophanyl-tRNA synthetase (trpS)
ORF02407 conserved hypothetical protein
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ORFxxxxx Annotation
ORF02408 ABC transporter, ATP-binding protein
ORF02409 ABC transporter, permease protein, putative
ORF02410 conserved hypothetical protein TIGR00246
ORF02411 serine protease
ORF02412 partitioning protein, ParB family
ORF02413 chromosomal replication initiator protein DnaA (dnaA)
ORF02415 DNA polymerase III, beta subunit (dnaN)
ORF02417 conserved hypothetical protein
ORF02419 conserved hypothetical GTP-binding protein
ORF02420 peptidyl-tRNA hydrolase (pth)
ORF02421 transcription-repair coupling factor (mfd)
ORF02423 S4 domain protein
ORF02424 cell division protein DivIC, putative
ORF02426 expressed protein of unknown function
ORF02427 MesJ/Ycf62 family protein
ORF02429 cell division protein FtsH (ftsH)

# Table 9: GBS genes shared with pneumoccocus

ORFxxxxx Ann tati n	
ODEO0047 - bear best best best best best best best best	
ORF00025 conserved hypothetical protein	urH)
ORF00029 acetyl xylan esterase, putative	
ORF00042 aldehyde-alcohol dehydrogenase (adhE)	
ORF00044 threonine synthase (thrC)	
ORF00044 threonine synthase (thrc) ORF00081 ribosomal protein L17 (rplQ)	
ORFO0000	
ORF00090 conserved hypothetical protein	
ORF00129 argininosuccinate synthase (argG)	
ORF00156 oligopeptide ABC transporter, substrate-binding protein, putative	
ORF00189 protease, putative	
ORF00194 thioredoxin family protein	
ORF00195 tRNA binding domain protein	
ORF00217 conserved domain protein	
ORF00218 PTS system, IIB component, putative	
ORF00220 transketolase, N-terminal subunit	
ORF00221 transketolase, C-terminal subunit	
ORF00223 oxidoreductase, putative	
ORF00282 acetyltransferase, GNAT family	<del></del>
ORF00290 IS1381, transposase OrfB	
ORF00291 IS1381, transposase OrfA	
ORF00293 conserved hypothetical protein	
ORF00301 membrane protein, putative	
ORF00343 ABC transporter, permease protein, putative	
ORF00344 conserved hypothetical protein	·
ORF00382 aspartate kinase family protein	
ORF00399 conserved hypothetical protein	100
ORF00439 cell wall surface anchor family protein	
ORF00447 cytidine/deoxycytidylate deaminase family protein	
ORF00450 5-formyltetrahydrofolate cyclo-ligase family protein	
ORF00480 transcriptional regulator, MerR family	
ORF00499 acetyltransferase, GNAT family	
ORF00504 magnesium transporter, CorA family	
ORF00521 VanZF domain protein	
ORF00612 IS1381, transposase OrfA	
ORF00613 IS1381, transposase OrfB	
ORF00690 transmembrane protein Vexp1 (vex1)	_
ORF00691 ABC transporter, ATP-binding protein Vexp2 (vex2)	
ORF00692 transmembrane protein Vexp3 (vex3)	
ORF00714 conserved hypothetical protein	
ORF00732 expressed cell wall surface anchor family protein, putative	
ORF00774 ABC transporter, ATP-binding protein	·
ORF00778 ABC transporter, ATP-binding protein	
ORF00778 conserved hypothetical protein	
ORF00790 beta-glucurőnidase	······
ORF00807 amina anid ARC territory	
ORF00807 amino acid ABC transporter, permease protein	
ORF00809 amino acid ABC transporter, amino acid-binding protein	
ORF00814 conserved hypothetical protein	
ORF00823 bacterial luciferase family protein	
ORF00840 riboflavin biosynthesis protein RibD (ribD)	
ORF00841 riboflavin synthase, alpha subunit (ribE)	
ORF00842 riboflavin biosynthesis protein RibA (ribA)	
ORF00843 riboflavin synthase, beta subunit (ribH)	
ORF00866 penicillin-binding protein 2b ORF00905 membrane protein, putative	

## Table 9: GBS g nes shared with pneumoc ocus

ORFxxxxx Ann tati n
ORF00910 major facilitator family protein
ORF00913 hydrolase, haloacid dehalogenase-like family
ORF00918 conserved hypothetical protein
ORF00945 conserved hypothetical protein
ORF00948 ABC transporter, ATP-binding protein
ORF00952 phosphomethylpyrimidine kinase (thiD)
ORF00953 hydroxyethylthiazole kinase (thiM)
ORF00954 thiamine-phosphate pyrophosphorylase (thiE)
ORF00961 GtrA family protein
ORF00967 1,4-alpha-glucan branching enzyme (glgB)
ORF00968 glucose-1-phosphate adenylyltransferase (glgC)
ORF00971 glycogen synthase (glgA)
ORF00985 acetyltransferase, GNAT family
ORF00990 magnesium transporter, CorA family, putative
ORF01022 nucleoside diphosphate kinase (ndk)
ORF01022 hdcleoside diphosphate kinase (hdk) ORF01031 nucleoside diphosphate kinase domain protein
ORF01085 conserved hypothetical protein
ORF01085 conserved hypothetical protein ORF01087 IS1381, transposase OrfA
ORF01087 IS1381, transposase OrfA ORF01088 IS1381, transposase OrfB
ORF01098 ABC transporter, permease protein, putative
ORF01100 sensor histidine kinase
ORF01102 ABC transporter, substrate-binding protein
ORF01127 protease, putative
ORF01135 iron compound ABC transporter, permease protein
ORF01136 iron compound ABC transporter, permease protein
ORF01185 aspartate-semialdehyde dehydrogenase (asd)
ORF01217 conserved hypothetical protein
ORF01218 conserved hypothetical protein
ORF01219 formate/nitrite transporter family protein
ORF01226 oxidoreductase, short chain dehydrogenase/reductase family, FRAMESHIFT
ORF01254 homoserine kinase (thrB)
ORF01255 homoserine dehydrogenase (hom)
ORF01264 transcriptional regulator, Cro/CI family
ORF01268 thiol peroxidase (psaD)
ORF01305 glycosyltransferase CpsJ(V) (cpsJ)
ORF01306 glycosyltransferase CpsO(V) (cpsO)
ORF01313 CpsD protein (cpsD)
ORF01314 cpsC protein (cpsC)
ORF01315 capsular polysaccharide biosynthesis protein CpsB (cpsB)
ORF01316 capsular polysaccharide biosynthesis protein CpsA (cpsA)
ORF01326 conserved hypothetical protein
ORF01333 alpha-acetolactate decarboxylase (budA)
ORF01334 acetolactate synthase, catabolic (ilvK)
ORF01337 MutT/nudix family protein
ORF01369 MATE efflux family protein
ORF01398 Tn5252, Orf 9 protein
ORF01399 Tn5252, Orf 10 protein
ORF01446 protease, putative
ORF01447 conserved hypothetical protein
ORF01449 conserved hypothetical protein
ORF01492 NADP-specific glutamate dehydrogenas (gdhA)
ORF01569 expressed cell wall surface anchor family protein
ORF01570 cell wall surface anchor family protein
IORE01574 polysaccharide biosynthesis protein
ORF01574 polysaccharide biosynthesis protein ORF01579 nucleotidyl transferase, putative

## Tabl 9: GBS genes shar d with pn umoccocus

ORFxxxxx Annotati n
ORF01580 polysaccharide biosynthesis protein, putative
ORF01612 conserved hypothetical protein
ORF01613 glycosyl transferase, group 1 family protein
ORF01617 conserved hypothetical protein
ORF01618 conserved hypothetical protein
ORF01621 glycosyl transferase, putative
ORF01622 glycosyl transferase, group 2 family protein
ORF01623 glycosyl transferase, family 8, degenerate
ORF01624 IS1381, transposase OrfB
ORF01625 IS1381, transposase OrfA
ORF01626 glycosyl transferase family 8
ORF01627 glycosyl transferase, family 8
ORF01628 conserved hypothetical protein
ORF01630 cell wall surface anchor family protein
ORF01635 protease, putative
ORF01643 aminopeptidase PepS (pepS)
ORF01702 peptidase, M20/M25/M40 family
ORF01731 IS1381, transposase OrfA
ORF01732 IS1381, transposase OrfB
ORF01740 tellurite resistance protein TehB (tehB)
ORF01747 methylated-DNAprotein-cysteine S-methyltransferase (ogt)
ORF01749 acetyltransferase, GNAT family
ORF01763 AcuB family protein
ORF01764 branched-chain amino acid ABC transporter, ATP-binding protein (livF)
ORF01765 branched-chain amino acid ABC transporter, ATP-binding protein (livG)
ORF01766 branched-chain amino acid ABC transporter, permease protein
ORF01767 branched-chain amino acid ABC transporter, permease protein (livH)
ORF01769 branched-chain amino acid ABC transporter, amino acid-binding protein
ORF01775 aminotransferase, class I
ORF01779 potassium uptake protein, Trk family
ORF01780 cation uptake protein, Trk family
ORF01824 cobalt transport family protein
ORF01826 conserved hypothetical protein
ORF01832 peptidase, M20/M25/M40 family
ORF01845 conserved hypothetical protein
ORF01848 transcriptional regulator, MerR family
ORF01853 isochorismatase family protein
ORF01859 membrane protein
ORF01875 oxidoreductase, aldo/keto reductase family
ORF01880 phospho-2-dehydro-3-deoxyheptonate aldolase
ORF01981 rRNA (guanine-N1-)-methyltransferase, putative
ORF02083 prophage LambdaSa2, DNA replication protein DnaC, putative
ORF02101 Na+/H+ exchanger family protein
ORF02107 membrane protein, putative
ORF02139 UDP-glucose 4-epimerase (galE)
ORF02143 lacX protein
ORF02162 conserved hypothetical protein
ORF02186 hemolysin precursor, putative
ORF02192 transcriptional regulator, MerR family
ORF02195 MutT/nudix family protein
ORF02228 IS1381, transposase OrfB
ORF02229 IS1381, transposase OrfA
ORF02233 conserved hypothetical protein
ORF02234 conserved hypothetical protein
ORF02276 5-methyltetrahydropteroyltriglutamatehomocysteine methyltransferase (metE)



## Tabl 9: GBS g n s shar d with pn um ccocus

ORFxxxxx Annotation
ORF02278 branched-chain amino acid transport protein AzIC, putativ
ORF02288 glycosyl transferase, family 8
ORF02289 glycosyl transferase, family 8
ORF02341 ribosomal protein L32 (rpmF)
ORF02343 conserved hypothetical protein
ORF02358 sensor histidine kinase
ORF02369 conserved hypothetical protein
ORF02384 LysM domain protein
ORF02428 hypoxanthine-guanine phosphoribosyltransferase (hpt)
ORF03011 ribosomal protein L33
ORF03014 ribosomal protein L33

# Table 10: GBS genes shar d with GAS

ORFXXXXX	Ann i	tation

ORFxxxx Ann tation
ORF00064 ribosomal protein S14, putative
ORF00095 D-alanyl-D-alanine carboxypeptidase family protein
ORF00096 N-acetylmuramoyl-L-alanine amidase, family 4 protein
ORF00110 conserved hypothetical protein
ORF00112 DNA repair protein RadA (radA)
ORF00124 permease, putative
ORF00148 glycosyl transferase, group 4 family protein
ORF00154 penicillin-binding protein 4, putative
ORF00157 oligopeptide ABC transporter, permease protein
ORF00206 oligopeptide ABC transporter, oligopeptide-binding protein
ORF00207 ollgopeptide ABC transporter, permease protein
ORF00208 oligopeptide ABC transporter, permease protein
ORF00209 peptide ABC transporter, ATP-binding protein
ORF00210 peptide ABC transporter, ATP-binding protein
ORF00216 IS1548, transposase
ORF00226 conserved hypothetical protein
ORF00232 conserved hypothetical protein
ORF00239 site-specific recombinase, phage integrase family
ORF00250 conserved hypothetical protein
ORF00251 conserved hypothetical protein
ORF00289 ABC transporter, ATP-binding protein
ORF00305 NADH oxidase, putative
ORF00317 cell division protein FtsL, putative
ORF00333 conserved hypothetical protein
ORF00383 hydrolase, haloacid dehalogenase-like family
ORF00430 expressed putative lipoprotein
ORF00431 transcriptional repressor CopY
ORF00434 membrane protein, putative
ORF00438 transcriptional regulator, Fur family
ORF00442 membrane protein, putative
QRF00445 bioY family protein
ORF00446 AtsA/ElaC family protein
ORF00468 expressed putative protease
ORF00469 glycosyl transferase, group 2 family protein
ORF00471 nrdl protein (nrdl)
ORF00473 expressed protein of unknown function
ORF00474 conserved hypothetical protein
ORF00507 conserved hypothetical protein
ORF00525 bioY family protein
ORF00528 thiolase
ORF00531 AMP-binding enzyme domain protein
ORF00548 YGGT family protein
ORF00565 exodeoxyribonuclease VII, small subunit (xseB)
ORF00565 exodeoxyribonuclease VII, small subunit (xseB) ORF00568 arginine repressor ArgR, putative
ORF00565 exodeoxyribonuclease VII, small subunit (xseB) ORF00568 arginine repressor ArgR, putative ORF00572 expressed putative lipase/acylhydrolase
ORF00565 exodeoxyribonuclease VII, small subunit (xseB) ORF00568 arginine repressor ArgR, putative ORF00572 expressed putative lipase/acylhydrolase ORF00573 conserved hypothetical protein
ORF00565 exodeoxyribonuclease VII, small subunit (xseB) ORF00568 arginine repressor ArgR, putative ORF00572 expressed putative lipase/acylhydrolase ORF00573 conserved hypothetical protein ORF00586 iron-sulfur cluster-binding protein, putative
ORF00565 exodeoxyribonuclease VII, small subunit (xseB) ORF00568 arginine repressor ArgR, putative ORF00572 expressed putative lipase/acylhydrolase ORF00573 conserved hypothetical protein ORF00586 iron-sulfur cluster-binding protein, putative ORF00592 oxidoreductase, short chain dehydrogenase/reductase family
ORF00565 exodeoxyribonuclease VII, small subunit (xseB) ORF00568 arginine repressor ArgR, putative ORF00572 expressed putative lipase/acylhydrolase ORF00573 conserved hypothetical protein ORF00586 iron-sulfur cluster-binding protein, putative ORF00592 oxidoreductase, short chain dehydrogenase/reductase family ORF00604 dipeptidase
ORF00565 exodeoxyribonuclease VII, small subunit (xseB) ORF00568 arginine repressor ArgR, putative ORF00572 expressed putative lipase/acylhydrolase ORF00573 conserved hypothetical protein ORF00586 iron-sulfur cluster-binding protein, putative ORF00592 oxidoreductase, short chain dehydrogenase/reductase family ORF00604 dipeptidase ORF00611 voltage-gated chloride channel family protein
ORF00565 exodeoxyribonuclease VII, small subunit (xseB) ORF00568 arginine repressor ArgR, putative ORF00572 expressed putative lipase/acylhydrolase ORF00573 conserved hypothetical protein ORF00586 iron-sulfur cluster-binding protein, putative ORF00592 oxidoreductase, short chain dehydrogenase/reductase family ORF00604 dipeptidase ORF00611 voltage-gated chloride channel family protein ORF00619 prophage LambdaSa1, repressor protein, putativ
ORF00565 exodeoxyribonuclease VII, small subunit (xseB) ORF00568 arginine repressor ArgR, putative ORF00572 expressed putative lipase/acylhydrolase ORF00573 conserved hypothetical protein ORF00586 iron-sulfur cluster-binding protein, putative ORF00592 oxidoreductase, short chain dehydrogenase/reductase family ORF00604 dipeptidase ORF00611 voltage-gated chloride channel family protein
ORF00565 exodeoxyribonuclease VII, small subunit (xseB) ORF00568 arginine repressor ArgR, putative ORF00572 expressed putative lipase/acylhydrolase ORF00573 conserved hypothetical protein ORF00586 iron-sulfur cluster-binding protein, putative ORF00592 oxidoreductase, short chain dehydrogenase/reductase family ORF00604 dipeptidase ORF00611 voltage-gated chloride channel family protein ORF00619 prophage LambdaSa1, repressor protein, putativ
ORF00565 exodeoxyribonuclease VII, small subunit (xseB) ORF00568 arginine repressor ArgR, putative ORF00572 expressed putative lipase/acylhydrolase ORF00573 conserved hypothetical protein ORF00586 iron-sulfur cluster-binding protein, putative ORF00592 oxidoreductase, short chain dehydrogenase/reductase family ORF00604 dipeptidase ORF00611 voltage-gated chloride channel family protein ORF00619 prophage LambdaSa1, repressor protein, putative ORF00622 conserved hypothetical protein ORF00627 prophag LambdaSa1, antirepressor, putative
ORF00565 exodeoxyribonuclease VII, small subunit (xseB) ORF00568 arginine repressor ArgR, putative ORF00572 expressed putative lipase/acylhydrolase ORF00573 conserved hypothetical protein ORF00586 iron-sulfur cluster-binding protein, putative ORF00592 oxidoreductase, short chain dehydrogenase/reductase family ORF00604 dipeptidase ORF00611 voltage-gated chloride channel family protein ORF00619 prophage LambdaSa1, repressor protein, putativ ORF00622 conserved hypothetical protein

# Table 10: GBS g nes shar d with GAS

ORFxxxxx Annotati n
ORF00654 conserved hypothetical protein
ORF00655 conserved hypothetical protein
ORF00656 conserved hypothetical protein
ORF00658 conserved hypothetical protein
ORF00659 conserved hypothetical protein
ORF00660 prophage LambdaSa1, structural protein, putative
ORF00662 conserved hypothetical protein
ORF00663 conserved hypothetical protein
ORF00664 conserved hypothetical protein
ORF00665 conserved hypothetical protein
ORF00666 prophage LambdaSa1, structural protein
ORF00668 conserved hypothetical protein
ORF00669 prophage LambdaSa1, pblA protein, internal deletion
ORF00667 prophage LambdaSa1, lysin, putative
ORF00679 conserved hypothetical protein
ORF00695 transposase OrfB, IS3 family, truncation
ORF00697 conserved hypothetical protein
ORF00707 conserved domain protein
ORF00713 acid phosphatase precursor, class B
ORF00720 transposase OrfB, IS3 family FRAMESHIFT
ORF00721 transposase OrfA, IS3 family
ORF00751 cylA protein (cylA)
ORF00755 cyll protein (cyll)
ORF00760 serine protease, subtilase family, putative POINT MUTATION
ORF00781 transcriptional regulator, LysR family
ORF00783 regulatory protein, putative
ORF00785 IS1548, transposase
ORF00786 regulatory protein, putative, truncation
ORF00787 D-lactate dehydrogenase (ldhA)
ORF00801 glycosyl transferase, group 1 family protein
ORF00805 conserved hypothetical protein
ORF00826 phage shock protein C, putative
ORF00833 conserved hypothetical protein
ORF00845 hydrolase, haloacid dehalogenase-like family
ORF00852 conserved hypothetical protein
ORF00853 expressed putative lipoprotein
ORF00857 IS1548, transposase
ORF00890 conserved hypothetical protein
ORF00902 conserved hypothetical protein
ORF00926 membrane protein, putative
ORF00927 membrane protein, putative
ORF00987 conserved hypothetical protein
ORF01009 expressed protein of unknown function
ORF01010 lipoyl-binding domain protein
ORF01011 oxidoreductase, putative
ORF01012 conserved hypothetical protein
ORF01024 expressed putative lipoprotein
ORF01061 signal peptidase I, putative
ORF01064 IS1548, transposase
ORF01084 glyoxylas family protein
ORF0104 SatD
ORF01104 SatD ORF01126 conserved hypothetical protein
ORF01191 conserved hypothetical protein
ORF01191 conserved hypothetical protein
ORF01193 glycine cleavage system H protein, putative
OKTO 1180 glydlie dieavage system 11 protein, putative

*'* 

## Table 10: GBS genes shared with GAS

ORFxxxxx Annotati n
ORF01194 bacterial luciferase family protein
ORF01195 oxidoreductase, FMN-binding
ORF01197 lipoate-protein ligase A family protein
ORF01202 IS861, transposase OrfA
ORF01223 drug resistance transporter, EmrB/QacA family, putative
ORF01224 conserved hypothetical protein
ORF01225 potassium uptake protein, putative
ORF01237 membrane protein, putative
ORF01249 dihydroneopterin aldolase (folB)
ORF01256 polysaccharide deacetylase family protein
ORF01273 transcriptional regulator, GntR family/potassioum uptake protein, TrkA family
ORF01280 conserved hypothetical protein
ORF01281 conserved hypothetical protein
ORF01289 lipoprotein, putative
ORF01291 conserved hypothetical protein
ORF01298 conserved hypothetical protein
ORF01318 conserved hypothetical protein
ORF01320 voltage-gated chloride channel family protein, putative
ORF01322 arsenate reductase (arsC)
ORF01340 dTDP-glucose 4,6-dehydratase (rfbB)
ORF01341 dTDP-4-dehydrorhamnose 3,5-epimerase
ORF01342 glucose-1-phosphate thymidylyltransferase (rfbA)
ORF01356 hypothetical protein
ORF01368 conserved hypothetical protein
ORF01374 ISSdy1, transposase OrfB
ORF01388 transposase OrfA, IS3 family
ORF01389 transposase OrfB, IS3 family, truncation
ORF01391 ISSdy1, transposase OrfB FRAMESHIFT
ORF01396 transcriptional regulator, Cro/Cl family
ORF01419 repressor protein, putative
ORF01461 amino acid permease
ORF01469 conserved hypothetical protein
ORF01483 sensor histidine kinase
ORF01485 GTP pyrophosphokinase family protein
ORF01490 5'-nucleotidase family protein
ORF01509 2-dehydropantoate 2-reductase, putative
ORF01510 regulatory protein, putative
ORF01522 carbamoyl-phosphate synthase, large subunit, putative
ORF01542 sulfatase
ORF01549 conserved hypothetical protein
ORF01554 iron compound ABC transporter, substrate-binding protein
ORF01557 conserved hypothetical protein
ORF01563 conserved hypothetical protein TIGR01212
ORF01583 glycosyltransferase, group 2 family protein
ORF01584 glycosyltransferase, group 2 family protein
ORF01585 glycosyltransferase, putative
ORF01586 dTDP-4-dehydrorhamnose reductase (rfbD)
ORF01593 conserved hypothetical protein
ORF01599 conserved hypothetical protein
ORF01600 glycerol-3-phosphate transporter, putative
ORF01639 conserved hypothetical protein
ORF01650 nitroreductas family protein
ORF01653 amino acid permease
ORF01665 transcriptional regulator, MutR family
ORF01683 MutT/nudix family protein
Ott 97000 Machinal terring protein

## Tabl 10: GBS genes shar d with GAS

ORFxxxxx Ann tation
ORF01686 67 kDa Myosin-crossreactive streptococcal antigen
ORF01688 peptide methionine sulfoxide reductase (msrA)
ORF01694 peptide ABC transporter, permease protein
ORF01704 conserved hypothetical protein
ORF01704 conserved hypothetical protein
ORF01703 13661, transposase OnA ORF01741 membrane protein, putative
ORF01741 memorane protein, putative
ORF01770 conserved hypothetical protein
ORF01772 IS1548, transposase
ORF01790 conserved hypothetical protein
ORF01794 conserved hypothetical protein
ORF01800 amino acid ABC transporter, substrate-binding protein
ORF01810 IS1548, transposase
ORF01827 sodium:dicarboxylate symporter family protein
ORF01877 immunogenic secreted protein, putative
ORF01913 transcriptional regulator, Cro/Cl family
ORF01928 membrane protein, putative
ORF01931 transporter, putative
ORF01932 transcriptional regulator, Crp/Fnr family
ORF01947 transcriptional regulator, merR family
ORF01970 acid phosphatase
ORF02002 amino acid ABC transporter, permease protein
ORF02028 perfringolysin O regulator protein (pfoR)
ORF02029 conserved hypothetical protein
ORF02031 expressed protein of unknown function
ORF02032 expressed protein of unknown function
ORF02035 deoxynucleoside kinase family protein
ORF02042 alkyl hydroperoxide reductase, subunit C (ahpC)
ORF02126 transcriptional regulator, MarR family
ORF02128 N-acetylmuramoyl-L-alanine amidase, family 4 protein
ORF02135 malate oxidoreductase
ORF02136 citrate carrier protein, CCS family
ORF02137 sensor histidine kinase family protein
ORF02138 response regulator
ORF02166 conserved hypothetical protein
ORF02169 PTS system, IIB component
ORF02170 PTS system, IIA component, putative
ORF02202 ABC transporter, ATP-binding protein
ORF02262 ABC transporter, ATP-binding protein
ORF02270 cAMP factor (cfb)
ORF02280 serine protease, subtilase family, putative
ORF02286 major facilitator family protein
ORF02292 preprotein translocase, SecE subunit, putative
ORF02295 Lyme disease proteins of unknown function, putative
ORF02298 Na+ dependent nucleoside transporter
ORF02301 transcriptional regulator, GntR family
ORF02313 virulence factor MviM, putative
ORF02316 membrane protein, putative
ORF02319 conserved hypothetical protein TIGR00250
ORF02328 transporter, putative
ORF02331 cold shock protein, CSD family
ORF02332 DNA mismatch repair protein HexA (h xA)
ORF02335 conserved hypothetical protein
ORF02372 conserved hypothetical protein
ORF02383 expressed putative lipoprotein
ORF02393 transporter, putativ
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## Table 10: GBS genes shared with GAS

ORFxxxxx Annotation	
ORF02398 transcriptional regulator, Crp/Fnr family	
ORF02399 conserved hypothetical protein	
ORF02401 acetyltransferase, GNAT family	
ORF02403 arginine/ornithine antiporter (arcD)	
ORF03002 conserved hypothetical protein, truncation	

## Table 11: GBS genes not shared with GAS r pneumococcus

100F 1 1.41
ORFxxxxx Ann tation
ORF00008 protease, putative
ORF00010 acyl carrier protein (acpP)
ORF00016 acetyltransferase, GNAT family
ORF00018 peptidase, M23/M37 family, putative secreted protein
ORF00035 membrane protein, putative
ORF00087 lipoprotein, putative
ORF00088 hypothetical protein
ORF00089 hypothetical protein
ORF00091 conserved hypothetical protein
ORF00117 ribose ABC transporter, periplasmic D-ribose-binding protein (rbsB)
ORF00118 ribose ABC transporter, permease protein (rbsC)
ORF00120 ribose ABC transporter protein RbsD (rbsD)
ORF00121 ribokinase (rbsK)
ORF00123 hypothetical protein
ORF00130 argininosuccinate lyase (argH)
ORF00137 conserved hypothetical protein
ORF00138 hypothetical protein
ORF00166 4-diphosphocytidyl-2C-methyl-D-erythritol kinase (ispE)
ORF00182 conserved domain protein
ORF00186 transcriptional regulator, Cro/Cl family
ORF00187 hypothetical protein
ORF00188 hypothetical protein
ORF00192 hypothetical protein
ORF00193 conserved hypothetical protein
ORF00196 conserved hypothetical protein
ORF00199 hydrolase, haloacid dehalogenase-like family
ORF00200 sensor histidine kinase, putative
ORF00201 response regulator
ORF00203 conserved hypothetical protein
ORF00204 membrane protein, putative
ORF00205 hypothetical protein
ORF00228 lipoprotein, putative
ORF00234 hypothetical protein
ORF00235 hypothetical protein
ORF00238 hypothetical protein
ORF00240 transcriptional regulator, Cro/CI family
ORF00241 hypothetical protein
ORF00242 conserved hypothetical protein
ORF00243 hypothetical protein
ORF00244 conserved domain protein
ORF00245 conserved hypothetical protein, fusion
ORF00246 replication initiation protein, putative
ORF00247 hypothetical protein
ORF00248 recombination protein
ORF00249 hypothetical protein
ORF00252 conserved hypothetical protein
ORF00253 hypothetical protein
ORF00254 hypothetical protein
ORF00255 hypothetical protein
ORF00255 hypothetical protein
ORF00257 hypothetical protein
ORF00258 hypothetical protein
ORF00259 hypothetical protein
ORF00260 hypothetical protein
ORF00272 expressed putative lipoprotein

# Tabl 11: GBS g nes not shared with GAS r pn umococcus

DRFxxxxx Ann tation DRF00273 hypothetical protein DRF00274 hypothetical protein DRF00275 hypothetical protein
DRF00274 hypothetical protein DRF00275 hypothetical protein
DRF00275 hypothetical protein
DRF00275 hypothetical protein
DRF00276 hypothetical protein
DRF00278 membrane protein, putative
DRF00279 transcriptional regulator, Cro/CI family
DRF00280 acetyltransferase, GNAT family
DRF00281 acetyltransferase, GNAT family
ORF00283 conserved hypothetical protein
ORF00284 RNA polymerase sigma factor, ECF subfamily
ORF00285 lipoprotein, putative
ORF00287 transcriptional regulator, TetR family
ORF00288 ABC transporter efflux protein, DrrB family, putative
ORF00292 hypothetical protein
ORF00294 expressed protein of unknown function
ORF00298 acyl carrier protein phosphodiesterase, putative
ORF00308 conserved hypothetical protein
ORF00324 conserved hypothetical protein
ORF00324 conserved hypothetical protein ORF00332 hypothetical protein
ORF00340 hypothetical protein
ORF00347 conserved hypothetical protein
ORF00384 hypothetical protein
ORF00402 membrane protein, putative
ORF00408 hypothetical protein
ORF00409 membrane protein, putative
ORF00414 conserved hypothetical protein
ORF00416 hypothetical protein
ORF00417 hypothetical protein
ORF00433 copper-transporter protein CopZ
ORF00448 hypothetical protein
ORF00466 conserved hypothetical protein
ORF00467 acetyltransferase, GNAT family
ORF00475 conserved domain protein
ORF00476 hypothetical protein
ORF00478 carboxymuconolactone decarboxylase family protein
ORF00479 conserved hypothetical protein
ORF00486 transcriptional regulator, AraC family
ORF00487 surface protein Rib
ORF00488 transposase, IS256 family, truncation
ORF00489 DNA-damage-inducible protein J, putative
ORF00490 hypothetical protein
ORF00491 lipoprotein, putative
ORF00493 bacteriophage L54a, integrase, truncation
ORF00497 conserved domain protein
ORF00503 oxidoreductase, Gfo/Idh/MocA family
ORF00506 transposase, IS256 family
ORF00510 bacteriocin transport accessory protein, putative
ORF00510 bacteriocin transport accessory protein, putative
ORF00526 biotin synthetase (bioB)
ORF00527 hypothetical protein
ORF00533 type IV prepilin peptidase-related protein
ORF00538 conserved hypothetical protein
ORF00556 hypothetical protein
ORF00563 xpressed protein of unknown function
ORF00575 hypothetical protein

## Tabl 11: GBS gen s not shared with GAS r pneum coccus

ORFxxxxx Ann tation
ORF00584 conserved hypothetical protein
ORF00585 fructose-1,6-bisphosphatase, putative
ORF00590 carboxymethylenebutenolidase-related protein
ORF00597 conserved hypothetical protein
ORF00598 inosine-uridine preferring nucleoside hydrolase
ORF00599 hypothetical protein
ORF00600 OsmC/Ohr family protein
ORF00608 adenosine deaminase, putative
ORF00610 chorismate mutase, putative
ORF00615 prophage LambdaSa1, site-specific recombinase, phage integrase family
ORF00617 conserved domain protein
ORF00618 hypothetical protein
ORF00620 hypothetical protein
ORF00621 conserved hypothetical protein
ORF00623 hypothetical protein
ORF00624 hypothetical protein
ORF00626 prophage LambdaSa1, transcriptional regulator, Cro/Cl family
ODECOCOO by a short of protein
ORF00628 hypothetical protein
ORF00630 hypothetical protein
ORF00632 hypothetical protein
ORF00633 conserved hypothetical protein
ORF00635 hypothetical protein
ORF00636 hypothetical protein
ORF00637 hypothetical protein
ORF00638 conserved hypothetical protein
ORF00639 conserved domain protein
ORF00641 prophage LambdaSa1, reverse transcriptase/maturase family protein
ORF00642 conserved hypothetical protein
ORF00643 conserved hypothetical protein
ORF00644 hypothetical protein
ORF00645 hypothetical protein
ORF00646 conserved hypothetical protein
ORF00647 hypothetical protein
ORF00649 hypothetical protein
ORF00650 hypothetical protein
ORF00652 conserved hypothetical protein
ORF00653 conserved hypothetical protein
ORF00657 conserved hypothetical protein, truncation
ORFO0007 Conserved hypothetical protein, indication
ORF00661 conserved hypothetical protein
ORF00667 conserved hypothetical protein
ORF00670 prophage LambdaSa1, minor structural protein, putative
ORF00671 prophage LambdaSa1, N-acetylmuramoyl-L-alanine amidase, family 4
ORF00672 prophage LambdaSa1, minor structural protein, putative
ORF00673 hypothetical protein
ORF00674 hypothetical protein
ORF00675 conserved hypothetical protein
ORF00676 conserved hypothetical protein
ORF00678 conserved hypothetical protein
ORF00681 conserved hypothetical protein
ORF00682 hypothetical protein
ORF00683 prophag LambdaSa1, site-specific recombinase, phage integrase family FRAMESHIFT
ORF00685 conserved hypothetical protein
ORF00689 conserved hypothetical protein, FRAMESHIFT
ORF00698 hypothetical protein
ORF00698 hypothetical protein ORF00703 phosphoserine phosphatase SerB (serB)
OKPUU/U3 phosphosenne phosphakase aera (sera)



# Tabl 11: GBS g nes not shared with GAS or pn um coccus

RFXXXX Annotation RF00712 hypothetical protein RF00712 hypothetical protein RF00713 hypothetical protein RF00723 hypothetical protein RF00725 transcriptional regulator, AraC family RF00726 capressed cell wall surface anchor family protein RF00726 expressed cell wall surface anchor family protein RF00726 expressed cell wall surface anchor family protein RF00737 conserved hypothetical protein, degenerate RF00737 conserved hypothetical protein RF00737 protein cell wall surface anchor family protein RF00737 protein cell protein RF00737 protein cell protein RF00738 hypothetical protein RF00740 hypothetical protein RF00741 hypothetical protein RF00742 hypothetical protein RF00747 cyll protein (cyll) RF00747 cyll protein (cyll) RF00747 cyll protein (cyll) RF00750 cyll protein (cyll) RF00750 cyll protein (cyll) RF00753 cyll protein (cyll) RF00753 cyll protein (cyll) RF00754 cyll protein (cyll) RF00756 cyll protein (cyll) RF00757 cyll protein (cyll) RF00757 cyll k protein (cyll) RF00759 putalive secreted protein RF00759 putalive secreted protein RF00769 cyll protein (cyll) RF00769 cyll protein (cyll) RF00769 cyll protein (cyll) RF00769 cyll protein (cyll) RF00769 protein (cyll) RF00769 cyll cyll protein (cyll) RF00769 cyll cyll protein (cyll) RF00769 cyll cyll cyll cyll cyll cyll cyll cyl	
IRF00712 hypothetical protein IRF00723 hypothetical protein IRF00726 transcriptional regulator, AraC family IRF00726 was pressed cell wall surface anchor family protein IRF007276 expressed cell wall surface anchor family protein IRF00728 expressed cell wall surface anchor family protein IRF00738 expressed cell wall surface anchor family protein IRF00738 expressed protein of unknown function IRF00738 hypothetical protein of unknown function IRF00738 hypothetical protein IRF00738 hypothetical protein IRF00740 hypothetical protein IRF00741 hypothetical protein IRF00741 hypothetical protein IRF00742 hypothetical protein IRF00742 hypothetical protein IRF00743 hypothetical protein IRF00743 hypothetical protein IRF00743 hypothetical protein AppC IRF00746 hypothetical protein hypothy	ORFxxxxx Annotation
INFO0773 cyll wall surface protein, interruption-N INFO0726 transcriptional regulator, AraC family INFO0726 transcriptional regulator, AraC family INFO0726 expressed cell wall surface anchor family protein INFO0736 expressed cell wall surface anchor family protein INFO0736 expressed protein of unknown function INFO0736 protein for info info info info info info info info	ORF00704 MutT/nudix family protein
IRF00723 hypothetical protein PRF00726 transcriptional regulator, AraC family PRF00727 expressed cell wall surface anchor family protein PRF00728 expressed cell wall surface anchor family protein PRF00738 expressed cell wall surface anchor family protein PRF00738 expressed protein of unknown function PRF00738 hypothetical protein of unknown function PRF00738 hypothetical protein PRF00749 hypothetical protein PRF00741 hypothetical protein PRF00741 hypothetical protein PRF00742 protein (cyll) PRF00742 explication protein expressed protein PRF00742 protein (cyll) PRF00749 explication protein explication PRF00749 explication protein explication PRF00749 explication protein explication PRF00750 explication protein explication PRF00750 explication explication PRF00750 explication explication PRF00750 explication explication PRF00756 explication explication PRF00756 explication explication PRF00756 expressed putative socreted protein PRF00756 expressed putative PRF007577 protein explication PRF00758 expressed putative PRF00758 expressed putative PRF00759 expressed putative PRF00759 expressed putative PRF00759 expressed putative PRF00759 membrane protein, putative PRF00759 membrane protein, putative PRF00759 D-mannonate exidoreductase exidence exidenc	ORF00712 hypothetical protein
NRF00726 transcriptional regulator, AraC family RRF00727 expressed cell wall surface anchor family protein NRF00736 expressed cell wall surface anchor family protein NRF00736 expressed protein of unknown function NRF00736 to sonserved hypothetical protein, degenerate NRF00738 hypothetical protein NRF00749 hypothetical protein NRF00741 hypothetical protein NRF00741 hypothetical protein NRF00742 lipoprotein, putative NRF00742 lipoprotein, putative NRF00742 protein for the cyll) NRF00743 expl protein (cyll) NRF00748 expl protein (cyll) NRF007582 cylls protein (cylls) NRF007584 cylls protein (cylls) NRF007585 cylls protein (cylls) NRF007585 cylls protein (cylls) NRF007586 cylls protein (cylls) NRF007586 hypothetical protein NRF00768 hypothetical protein NRF00768 expressed putative secreted protein NRF00768 cypressed putative secreted protein NRF00768 cypressed putative secreted protein NRF00768 conserved domain protein NRF00769 permease, putative NRF00779 potentical protein NRF00779 potentical protein, putative NRF00779 bod family protein, putative NRF00779 hypothetical protein NRF00798 hydrolase, haloacid dehalogenase-like family NRF00798 hydrolase, haloacid dehalogenase-like family NRF00798 hypothetical protein potein potein NRF00888 hyp	ORF00718 cell wall surface protein, interruption-N
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ORF00779 membrane protein, putative ORF00788 sodium:galactoside symporter family protein, putative ORF00791 transcriptional regulator, GntR family ORF00793 Glucuronate isomerase (uxaC) ORF00794 mannonate dehydratase (uxuA) ORF00795 D-mannonate oxidoreductase ORF00796 hydrolase, haloacid dehalogenase-like family ORF00797 glycosyl hydrolase, family 3 ORF00806 conserved hypothetical protein ORF00822 ABC transporter, ATP-binding protein ORF00827 hypothetical protein ORF00834 conserved hypothetical protein ORF00838 membrane protein, putative ORF00839 Mn2+/Fe2+ transporter, NRAMP family ORF00848 conserved domain protein ORF00872 cell wall surface anchor family protein ORF00874 conserved hypothetical protein ORF00878 ABC transporter, permease protein ORF00879 YaeC family protein, putative ORF00888 hydrolase, haloacid dehalogenase-like family ORF00891 conserved domain protein	
ORF00788 sodium:galactoside symporter family protein, putative ORF00791 transcriptional regulator, GntR family ORF00793 Glucuronate isomerase (uxaC) ORF00794 mannonate dehydratase (uxuA) ORF00795 D-mannonate oxidoreductase ORF00796 hydrolase, haloacid dehalogenase-like family ORF00797 glycosyl hydrolase, family 3 ORF00806 conserved hypothetical protein ORF00822 ABC transporter, ATP-binding protein ORF00827 hypothetical protein ORF00834 conserved hypothetical protein ORF00838 membrane protein, putative ORF00839 Mn2+/Fe2+ transporter, NRAMP family ORF00848 conserved domain protein ORF00872 cell wall surface anchor family protein ORF00874 conserved hypothetical protein ORF00878 ABC transporter, permease protein ORF00879 YaeC family protein, putative ORF00888 hydrolase, haloacid dehalogenase-like family ORF00891 conserved domain protein	ORF00779 membrane protein, putative
ORF00791 transcriptional regulator, GntR family ORF00793 Glucuronate isomerase (uxaC) ORF00794 mannonate dehydratase (uxuA) ORF00795 D-mannonate oxidoreductase ORF00796 hydrolase, haloacid dehalogenase-like family ORF00797 glycosyl hydrolase, family 3 ORF00806 conserved hypothetical protein ORF00822 ABC transporter, ATP-binding protein ORF00827 hypothetical protein ORF00834 conserved hypothetical protein ORF00838 membrane protein, putative ORF00839 Mn2+/Fe2+ transporter, NRAMP family ORF00848 conserved domain protein ORF00872 cell wall surface anchor family protein ORF00874 conserved hypothetical protein ORF00878 ABC transporter, permease protein ORF00879 YaeC family protein, putative ORF00888 hydrolase, haloacid dehalogenase-like family ORF00891 conserved domain protein	ORF00798 sodium:galactoside sympotter family protein, putative
ORF00793 Glucuronate isomerase (uxuC) ORF00794 mannonate dehydratase (uxuA) ORF00795 D-mannonate oxidoreductase ORF00796 hydrolase, haloacid dehalogenase-like family ORF00797 glycosyl hydrolase, family 3 ORF00806 conserved hypothetical protein ORF00822 ABC transporter, ATP-binding protein ORF00827 hypothetical protein ORF00834 conserved hypothetical protein ORF00838 membrane protein, putative ORF00839 Mn2+/Fe2+ transporter, NRAMP family ORF00848 conserved domain protein ORF00872 cell wall surface anchor family protein ORF00874 conserved hypothetical protein ORF00878 ABC transporter, permease protein ORF00879 YaeC family protein, putative ORF00888 hydrolase, haloacid dehalogenase-like family ORF00891 conserved domain protein	ORF00701 transcriptional regulator. GotR family
ORF00794 mannonate dehydratase (uxuA) ORF00795 D-mannonate oxidoreductase ORF00796 hydrolase, haloacid dehalogenase-like family ORF0080797 glycosyl hydrolase, family 3 ORF00806 conserved hypothetical protein ORF00822 ABC transporter, ATP-binding protein ORF00827 hypothetical protein ORF00834 conserved hypothetical protein ORF00838 membrane protein, putative ORF00839 Mn2+/Fe2+ transporter, NRAMP family ORF00848 conserved domain protein ORF00872 cell wall surface anchor family protein ORF00874 conserved hypothetical protein ORF00878 ABC transporter, permease protein ORF00879 YaeC family protein, putative ORF00888 hydrolase, haloacid dehalogenase-like family ORF00891 conserved domain protein	ORF00793 (Italiscriptional regulator, Onto Tarinin)
ORF00795 D-mannonate oxidoreductase ORF00796 hydrolase, haloacid dehalogenase-like family ORF00806 conserved hypothetical protein ORF00822 ABC transporter, ATP-binding protein ORF00827 hypothetical protein ORF00834 conserved hypothetical protein ORF00838 membrane protein, putative ORF00839 Mn2+/Fe2+ transporter, NRAMP family ORF00848 conserved domain protein ORF00872 cell wall surface anchor family protein ORF00874 conserved hypothetical protein ORF00878 ABC transporter, permease protein ORF00879 YaeC family protein, putative ORF00888 hydrolase, haloacid dehalogenase-like family ORF00891 conserved domain protein	
ORF00796 hydrolase, haloacid dehalogenase-like family ORF00797 glycosyl hydrolase, family 3 ORF00806 conserved hypothetical protein ORF00822 ABC transporter, ATP-binding protein ORF00827 hypothetical protein ORF00834 conserved hypothetical protein ORF00838 membrane protein, putative ORF00839 Mn2+/Fe2+ transporter, NRAMP family ORF00848 conserved domain protein ORF00872 cell wall surface anchor family protein ORF00874 conserved hypothetical protein ORF00878 ABC transporter, permease protein ORF00879 YaeC family protein, putative ORF00888 hydrolase, haloacid dehalogenase-like family ORF00891 conserved domain protein	
ORF00806 conserved hypothetical protein ORF00822 ABC transporter, ATP-binding protein ORF00827 hypothetical protein ORF00834 conserved hypothetical protein ORF00838 membrane protein, putative ORF00839 Mn2+/Fe2+ transporter, NRAMP family ORF00848 conserved domain protein ORF00872 cell wall surface anchor family protein ORF00874 conserved hypothetical protein ORF00878 ABC transporter, permease protein ORF00879 YaeC family protein, putative ORF00888 hydrolase, haloacid dehalogenase-like family ORF00891 conserved domain protein	
ORF00806 conserved hypothetical protein ORF00822 ABC transporter, ATP-binding protein ORF00827 hypothetical protein ORF00834 conserved hypothetical protein ORF00838 membrane protein, putative ORF00839 Mn2+/Fe2+ transporter, NRAMP family ORF00848 conserved domain protein ORF00872 cell wall surface anchor family protein ORF00874 conserved hypothetical protein ORF00878 ABC transporter, permease protein ORF00879 YaeC family protein, putative ORF00888 hydrolase, haloacid dehalogenase-like family ORF00891 conserved domain protein	ORF00/96 hydroiase, naioacid denalogenase-like lanniny
ORF00822 ABC transporter, ATP-binding protein ORF00827 hypothetical protein ORF00834 conserved hypothetical protein ORF00838 membrane protein, putative ORF00839 Mn2+/Fe2+ transporter, NRAMP family ORF00848 conserved domain protein ORF00872 cell wall surface anchor family protein ORF00874 conserved hypothetical protein ORF00878 ABC transporter, permease protein ORF00879 YaeC family protein, putative ORF00888 hydrolase, haloacid dehalogenase-like family ORF00891 conserved domain protein	ORF00/9/ glycosyl nydrolase, lamily 3
ORF00827 hypothetical protein ORF00834 conserved hypothetical protein ORF00838 membrane protein, putative ORF00839 Mn2+/Fe2+ transporter, NRAMP family ORF00848 conserved domain protein ORF00872 cell wall surface anchor family protein ORF00874 conserved hypothetical protein ORF00878 ABC transporter, permease protein ORF00879 YaeC family protein, putative ORF00888 hydrolase, haloacid dehalogenase-like family ORF00891 conserved domain protein	ORF00806 ADC Assessed hypothetical protein
ORF00834 conserved hypothetical protein ORF00838 membrane protein, putative ORF00839 Mn2+/Fe2+ transporter, NRAMP family ORF00848 conserved domain protein ORF00872 cell wall surface anchor family protein ORF00874 conserved hypothetical protein ORF00878 ABC transporter, permease protein ORF00879 YaeC family protein, putative ORF00888 hydrolase, haloacid dehalogenase-like family ORF00891 conserved domain protein	
ORF00838 membrane protein, putative ORF00839 Mn2+/Fe2+ transporter, NRAMP family ORF00848 conserved domain protein ORF00872 cell wall surface anchor family protein ORF00874 conserved hypothetical protein ORF00878 ABC transporter, permease protein ORF00879 YaeC family protein, putative ORF00888 hydrolase, haloacid dehalogenase-like family ORF00891 conserved domain protein	ORF00827 hypothetical protein
ORF00839 Mn2+/Fe2+ transporter, NRAMP family ORF00848 conserved domain protein ORF00872 cell wall surface anchor family protein ORF00874 conserved hypothetical protein ORF00878 ABC transporter, permease protein ORF00879 YaeC family protein, putative ORF00888 hydrolase, haloacid dehalogenase-like family ORF00891 conserved domain protein	
ORF00848 conserved domain protein ORF00872 cell wall surface anchor family protein ORF00874 conserved hypothetical protein ORF00878 ABC transporter, permease protein ORF00879 YaeC family protein, putative ORF00888 hydrolase, haloacid dehalogenase-like family ORF00891 conserved domain protein	
ORF00872 cell wall surface anchor family protein ORF00874 conserved hypothetical protein ORF00878 ABC transporter, permease protein ORF00879 YaeC family protein, putative ORF00888 hydrolase, haloacid dehalogenase-like family ORF00891 conserved domain protein	ORF00839 Mn2+/Fe2+ transporter, NKAWP tamily
ORF00874 conserved hypothetical protein ORF00878 ABC transporter, permease protein ORF00879 YaeC family protein, putative ORF00888 hydrolase, haloacid dehalogenase-like family ORF00891 conserved domain protein	
ORF00878 ABC transporter, permease protein ORF00879 YaeC family protein, putative ORF00888 hydrolase, haloacid dehalogenase-like family ORF00891 conserved domain protein	
ORF00879 YaeC family protein, putative ORF00888 hydrolase, haloacid dehalogenase-like family ORF00891 conserved domain protein	
ORF00888 hydrolase, haloacid dehalogenase-like family ORF00891 conserved domain protein	ORF008/8 ABC transporter, permease protein
ORF00891 conserved domain protein	ORF00879 YaeC family protein, putative
IORF00898 conserved hypothetical protein	
ORF00900 permease, GntP family	ORF00900 permease, GntP family
ORF00903 transcriptional regulator, MarR family	ORF00903 transcriptional regulator, MarR family
ORF00907 glutathione S-transferase family protein	ORF00907 glutathione S-transferase family protein



ORFxxxxx Annotati n	
ORF00909 hypothetical protein	
ORF00921 membrane protein, putative	
ORF00922 glycosyl transferase, family 8	
ORF00923 hypothetical protein	
ORF00924 conserved hypothetical protein	
ORF00939 conserved hypothetical protein	
ORF00942 expressed putative secreted protein	
ORF00943 hypothetical protein	
ORF00944 hypothetical protein	
ORF00946 conserved hypothetical protein	
ORF00950 hypothetical protein	
ORF00951 transcriptional regulator, TenA family	
ORF00972 ATP synthase F0, C subunit (atpE)	
ORF00980 conserved hypothetical protein	
ORF00982 conserved hypothetical protein	
ORF01003 conserved hypothetical protein	
ORF01004 conserved hypothetical protein	
ORF01013 hypothetical protein	
ORF01014 hypothetical protein	
ON 010 13 hypothetical protein	
ORF01016 hypothetical protein	
ORF01018 hypothetical protein	
ORF01019 hypothetical protein	`
ORF01021 hypothetical protein	
ORF01025 HD domain protein	
ORF01026 acetyltransferase, GNAT family	
ORF01032 chloramphenicol acetyltransferase (cat)	
ORF01034 Tn916, transposase	
ORF01035 Tn916, excisionase	
ORF01037 Tn916, hypothetical protein	
ORF01038 Tn916, hypothetical protein	
ORF01039 Tn916, transcriptional regulator, putative	
ORF01041 Tn916, hypothetical protein	
ORF01042 Tn916, NLP/P60 family protein	
ORF01044 membrane protein, putative FRAMESHIFT	
ORF01048 Tn916, hypothetical protein	
ORF01049 Tn916, hypothetical protein	
ORF01050 Tn916, hypothetical protein	
ORF01051 Tn916, transcriptional regulator, putative ORF01052 Tn916, FtsK/SpolIIE family protein	
ORF01053 Tn916, hypothetical protein	
ORF01054 Tn916, hypothetical protein	
ORF01062 hypothetical protein	
ORF01086 Na+/H+ exchanger family protein	
ORF01092 acetyltransferase, GNAT family	·
ORF01096 nisin-resistance protein, putative	
ORF01103 conserved hypothetical protein	
ORF01124 acetyltransferase, GNAT family	
ORF01133 iron-compound ABC transporter, iron-compound-binding protein	
ORF01140 conserved hypothetical protein	
ORF01142 carbon starvation protein CstA, putative	
ORF01143 response regulator	
ORF01144 sensor histidine kinase, putative	
ORF01145 lipoprotein, putative	
ORF01146 conserved hypothetical protein, FRAMESHIFT	
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Tabl 11: GBS g n s not shared with GAS r pneumococcus

ORFxxxxx Annotation
ORF01148 lipoprotein, putative
ORF01149 hypothetical protein
ORF01150 hypothetical protein
ORF01151 hypothetical protein
ORF01152 lipoprotein, putative
ORF01153 hypothetical protein
ORF01157 conserved hypothetical protein
ORF01158 hypothetical protein
ORF01159 hypothetical protein
ORF01160 expressed protein of unknown function FRAMESHIFT
ORF01161 expressed conserved domain protein
ORF01162 conserved hypothetical protein
ORF01164 FtsK/SpollIE family protein FRAMESHIFT
ORF01166 hypothetical protein
ORF01167 conserved hypothetical protein
ORF01168 conserved hypothetical protein
ORF01169 hypothetical protein
ORF01172 phage infection protein, putative
ORF01173 conserved hypothetical protein
ORF01174 conserved domain protein
ORF01175 hypothetical protein
ORF01182 membrane protein, putative
ORF01186 cell wall surface anchor family protein, putative
ORF01187 hypothetical protein
ORF01204 hypothetical protein
ORF01215 hypothetical protein
ORF01241 transcriptional regulator, AraC family, putative
ORF01253 rarD protein (rarD)
ORF01257 transporter, BCCT family protein
ORF01258 hypothetical protein
ORF01261 expressed protein of unknown function
ORF01262 conserved hypothetical protein, FRAMESHIFT
ORF01263 hypothetical protein
ORF01265 hypothetical protein
ORF01266 hypothetical protein
ORF01269 conserved hypothetical protein
ORF01272 conserved hypothetical protein
ORF01277 conserved hypothetical protein
ORF01287 conserved hypothetical protein
ORF01288 membrane protein, putative
ORF01299 CMP-N-acetylneuraminic acid synthetase NeuA (neuA)
ORF01300 neuD protein (neuD)
ORF01301 UDP-N-acetylglucosamine-2-epimerase NeuC (neuC)
ORF01302 N-acetyl neuramic acid synthetase NeuB (neuB)
ORF01303 polysaccharide biosynthesis protein CpsL (cpsL)
ORF01304 polysaccharide biosynthesis protein CpsK(V) (cpsK)
ORF01307 glycosyltransferase CpsN(V) (cpsN)
ORF01308 polysaccharide biosynthesis protein CpsM(V) (cpsM)
ORF01309 polysaccharide biosynthesis protein cpsH(V) (cpsH)
ORF01310 glycosyltransferase CpsG(V) (cpsG)
ORF01311 polysaccharide biosynthesis protein CpsF (cpsF)
ORF01312 glycosyltransferase CpsE (cpsE)
ORF01348 conserved domain protein
ORF01349 hypothetical protein
ORF01370 conserved hypothetical protein

## Tabl 11: GBS gen s not shared with GAS r pneumococcus

ORFxxxxx Annotati n
ORF01371 conserv d hypothetical protein
ORF01372 expressed protein of unknown function
ORF01373 ISSdy1, transposase OrfA
ORF01375 conserved hypothetical protein
ORF01379 transposase OrfB, IS3 family, truncation
ORF01382 GBSi1, group II intron, maturase
ORF01384 hypothetical protein
ORF01385 hypothetical protein
ORF01386 conserved hypothetical protein
ORF01387 conserved hypothetical protein, truncation
ORF01390 ISSdy1, transposase OrfA FRAMESHIFT
ORF01392 hypothetical protein
ORF01393 hypothetical protein
ORF01394 site-specific recombinase, phage integrase family
ORF01395 conserved hypothetical protein
ORF01401 transposase, ISL3 family
ORF01401 transposase, ISLS family ORF01404 mercuric resistance operon regulatory protein MerR (merR)
ORF01404 mercunc resistance operon regulatory protein (merk)  ORF01408 cadmium efflux system accessory protein (CadC)
ORF01409 conserved hypothetical protein
ORF01410 hypothetical protein
ORF01417 hypothetical protein
ORF01418 hypothetical protein
ORF01420 hypothetical protein
ORF01421 ImpB/MucB/SamB family protein
ORF01423 conserved hypothetical protein
ORF01424 conserved hypothetical protein
ORF01425 conserved hypothetical protein
ORF01426 conserved hypothetical protein
ORF01427 hypothetical protein
ORF01428 conserved hypothetical protein
ORF01430 hypothetical protein
ORF01431 hypothetical protein
ORF01432 conserved domain protein
ORF01433 SNF2 family protein
ORF01434 hypothetical protein
ORF01435 calcium-binding protein, putative
ORF01436 agglutinin receptor (ssp-5)
ORF01437 abortive infection protein AbiGI (abiGI)
ORF01438 abortive infection protein AbiGII (abiGII)
ORF01439 conserved hypothetical protein
ORF01440 expressed protein of unknown function
ORF01441 conserved hypothetical protein, degenerate
ORF01441 conserved hypothetical protein, degenerate  ORF01442 membrane protein, putative
ORF01443 hypothetical protein
ORF01444 Tn5252, Orf 21 protein, internal deletion
ORF01445 hypothetical protein
ORF01450 conserved hypothetical protein
ORF01452 hypothetical protein
ORF01454 conserved hypothetical protein
ORF01459 hypothetical protein
ORF01460 homocysteine S-methyltransferase MmuM, putative
ORF01463 hypothetical protein
ORF01464 hypothetical protein
ORF01465 hypothetical protein
ORF01466 transcriptional regulator, TetR family
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ODE Annuatation
ORFxxxxx Annotation
ORF01477 glutathione S-transferase family protein, putative
ORF01478 conserved domain protein
ORF01486 hypothetical protein
ORF01488 R5 protein
ORF01489 transcriptional regulator, MarR family, putative
ORF01494 membrane protein, putative
ORF01497 acetyltransferase, GNAT family
ORF01502 hypothetical protein
ORF01503 conserved hypothetical protein
ORF01508 surface antigen-related protein
ORF01535 conserved hypothetical protein
ORF01547 conserved hypothetical protein
ORF01566 expressed cell wall surface anchor family protein
ORF01572 glycosyltransferase, group 1 family protein
ORF01573 glycosyltransferase, group 2 family protein
ORF01575 membrane protein, putative
ORF01576 glycosyltransferase, group 2 family protein
ORF01577 glycosyltransferase, group 2 family protein
ORF01578 nucleotide sugar dehydratase, putative
ORF01581 lipoprotein, putative
ORF01582 conserved hypothetical protein
ORF01596 ammonium transporter family protein
ORF01597 conserved hypothetical protein
ORF01601 hypothetical protein
ORF01608 proton/peptide symporter family protein
ORF01611 hypothetical protein
ORF01615 conserved domain protein
ORF01638 conserved hypothetical protein
ORF01641 conserved hypothetical protein
ORF01645 cell wall surface anchor family protein
ORF01660 membrane protein, putative
ORF01661 ABC transporter, ATP binding protein
ORF01666 hypothetical protein
ORF01667 hypothetical protein
ORF01670 hypothetical protein
ORF01672 protease, putative, POINT MUTATION
ORF01673 hypothetical protein
ORF01674 hypothetical protein
ORF01675 hypothetical protein
ORF01680 tetracenomycin polyketide synthesis O-methyltransferase TcmP, putative
ORF01681 hypothetical protein
ORF01682 hypothetical protein
ORF01684 hypothetical protein
ORF01692 peptide ABC transporter, ATP-binding protein
ORF01695 peptide ABC transporter, permease protein
ORF01696 peptide ABC transporter, peptide-binding protein
ORF01699 transposase, IS30 family, putative
ORF01700 transporter, major facilitator family
ORF01703 transcriptional regulator, LysR family
ORF01715 conserved hypothetical protein
ORF01719 hypothetical protein
ORF01720 conserved hypothetical protein
ORF01721 glyoxalase family protein
ORF01727 conserved hypothetical protein
ORF01729 acetyltransferase, GNAT family
10.4 o 1720 decly administrate, Grave landy



# Tabl 11: GBS g nes n t shared with GAS or pneumococcus

ORFxxxxx Annotation
ORF01730 glycosyl transferase, group 2 family protein
ORF01733 hypothetical protein
ORF01734 conserved hypothetical protein
ORF01735 hypothetical protein
ORF01736 hypothetical protein
ORF01737 hypothetical protein
ORF01742 hypothetical protein
ORF01743 PTS system component, putative
ORF01744 conserved hypothetical protein
ORF01748 D-isomer specific 2-hydroxyacid dehydrogenase family protein
ORF01753 conserved hypothetical protein
ORF01754 hypothetical protein
ORF01761 transposase, IS30 family, putative, truncation
ORF01778 amino acid permease, putative
ORF01807 hypothetical protein
ORF01836 hypothetical protein
ORF01838 hypothetical protein
ORF01839 dihydroxyacetone kinase family protein
ORF01840 transcriptional regulator, TetR family, putative
ORF01842 hypothetical protein
ORF01843 dihydroxyacetone kinase family protein
ORF01844 dihydroxyacetone kinase family protein
ORF01847 conserved hypothetical protein
ORF01850 hypothetical protein
ORF01863 pyruvate phosphate dikinase (ppdK)
ORF01864 expressed protein of unknown function
ORF01865 CBS domain protein
ORF01866 3-hydroxyacyl-CoA dehydrogenase family protein, putative secreted protein
ORF01892 hypothetical protein
ORF01893 hypothetical protein
ORF01894 conserved hypothetical protein
ORF01895 hypothetical protein
ORF01896 hypothetical protein
ORF01897 hypothetical protein
ORF01898 hypothetical protein
ORF01899 hypothetical protein
ORF01903 conserved hypothetical protein
ORF01904 drug resistance transporter, EmrB/QacA family
ORF01905 hypothetical protein
ORF01922 conserved hypothetical protein
ORF01925 FMN-binding protein
ORF01934 hypothetical protein
ORF01936 polyprenyl synthetase family protein
ORF01939 cytochrome d ubiquinol oxidase, subunit II (cydB)
ORF01940 cytochrome d oxidase, subunit I (cydA)
ORF01941 pyridine nucleotide-disulphide oxidoreductase family protein
ORF01942 prenyltransferase, UbiA family
ORF01943 hypothetical protein
ORF01944 hypothetical protein
ORF01946 cyclopropane-fatty-acyl-phospholipid synthase (cfa)
ORF01951 conserved hypothetical protein
ORF01953 hypothetical protein
ORF01954 conserved hypothetical protein
ORF01984 hypothetical protein
ORF01988 hypothetical protein
On orado hypothetical protein

# Tabl 11: GBS g nes not shared with GAS r pneumococcus

ODFAmpérélon	
ORFxxxxx Annotation	
ORF01989 hypothetical protein	
ORF01990 hypothetical protein	
ORF01991 hypothetical protein	
ORF02000 membrane protein, putative	
ORF02001 transposase, IS30 family, putative	
ORF02005 hypothetical protein	
ORF02006 xylulose-5-phosphate/fructose-6-phosphate phosphoketolase (xfp)	
ORF02009 conserved hypothetical protein	
ORF02010 carbohydrate kinase, FGGY family	
ORF02011 hypothetical protein	
ORF02012 PTS system component, putative	
ORF02015 glyoxylate reductase, NADH-dependent	
ORF02016 hypothetical protein	
ORF02025 hypothetical protein	
ORF02026 hypothetical protein	
ORF02030 glutamate-cysteine ligase-related protein	
ORF02036 phosphinothricin N-acetyltransferase (pat)	
ORF02039 conserved hypothetical protein	
ORF02044 conserved hypothetical protein	
ORF02045 conserved hypothetical protein	
ORF02046 prophage LambdaSa2, lysin, putative	
ORF02047 prophage LambdaSa2, holin, putative	
ORF02048 conserved hypothetical protein	
ORF02049 hypothetical protein	
ORF02050 conserved domain protein	
ORF02051 prophage LambdaSa2, PbIB, putative	
ORF02053 conserved hypothetical protein	
ORF02056 conserved hypothetical protein	
ORF02057 hypothetical protein	
ORF02058 hypothetical protein	
ORF02059 conserved hypothetical protein	
ORF02060 conserved hypothetical protein	
ORF02060 Conserved hypothetical protein	
ORF02062 hypothetical protein	
ORF02063 conserved domain protein	
ORF02064 conserved domain protein	
ORF02066 prophage LambdaSa2, protease, putative	
ORF02067 conserved hypothetical protein	
ORF02068 prophage LambdaSa2, terminase large subunit, putative	
ORF02069 hypothetical protein	
ORF02070 hypothetical protein	
ORF02071 prophage LambdaSa2, site-specific recombinase, phage integrase family	<del></del>
ORF02072 conserved hypothetical protein	
ORF02073 prophage LambdaSa2, transcriptional regulator, Cro/CI family	<del></del>
ORF02075 hypothetical protein	
ORF02077 hypothetical protein	
ORF02078 conserved hypothetical protein	
ORF02079 conserved hypothetical protein	
ORF02080 conserved hypothetical protein	
ORF02081 hypothetical protein	
ORF02084 prophage LambdaSa2, bacteriophage replication protein/hypothetical protein,	
truncation/fusion	
ORF02085 hypothetical protein	
ORF02087 hypothetical protein	
ORF02088 conserved hypothetical protein	
Otti v2000 conton roa riypourousus present	

# Tabl 11: GBS g n s not shared with GAS r pneum c ccus

ORFxxxxx Annotation
ORF02089 prophage LambdaSa2, HNH endonuclease family protein
ORF02090 prophage LambdaSa2, antirepressor protein, putative
ORF02091 conserved domain protein
ORF02092 hypothetical protein
ORF02093 hypothetical protein
ORF02094 hypothetical protein
ORF02095 prophage LambdaSa2, repressor protein, putative
ORF02097 hypothetical protein
ORF02098 prophage LambdaSa2, site-specific recombinase, phage integrase family
ORF02100 hypothetical protein
ORF02102 hypothetical protein
ORF02103 microcin immunity protein MccF, putative
ORF02105 oxidoreductase, Gfo/Idh/MocA family
ORF02108 hypothetical protein
ORF02109 Cyclic nucleotide-binding domain protein
ORF02119 hypothetical protein
ORF02124 hypothetical protein
ORF02125 nitroreductase family protein
ORF02134 bacteriocin transport accessory protein, putative
ORF02148 neuraminidase-related protein
ORF02160 2`,3`-cyclic-nucleotide 2`-phosphodiesterase (cpdB)
ORF02163 conserved hypothetical protein
ORF02171 membrane protein, putative
ORF02172 hypothetical protein
ORF02173 membrane protein, putative
ORF02175 conserved hypothetical protein, truncation
ORF02181 phosphate transport system regulatory protein PhoU, putative
ORF02187 hypothetical protein
ORF02190 conserved hypothetical protein
ORF02191 hypothetical protein
ORF02194 acetyltransferase, GNAT family
ORF02196 hypothetical protein
ORF02198 acetyltransferase, GNAT family
ORF02201 membrane protein, putative
ORF02203 hypothetical protein
ORF02205 transcriptional regulator, Cro/CI family
ORF02206 conserved hypothetical protein
ORF02207 conserved hypothetical protein TIGR00730
ORF02208 hypothetical protein
ORF02209 site-specific recombinase, phage integrase family
ORF02210 conserved hypothetical protein
ORF02211 conserved hypothetical protein
ORF02211 Conserved hypothetical protein
ORF02213 hypothetical protein
ORF02214 transcriptional regulator, Cro/CI family
ORF02215 expressed protein of unknown function
ORF02216 site-specific recombinase, phage integrase family
ORF02217 conserved hypothetical protein
ORF02219 hypothetical protein
ORF02221 cell wall anchor protein-related protein
ORF02223 hypothetical protein
ORF02224 hypothetical protein
ORF02225 hypothetical protein
ORF02226 membrane protein, putative
ORF02227 conjugal transfer protein, interruption-C

## Tabl 11: GBS gen s not shared with GAS r pneumoc ccus

ORFxxxxx Annotation
ORF02230 conserved hypothetical protein
ORF02231 conserved hypothetical protein
ORF02232 conserved hypothetical protein
ORF02235 hypothetical protein
ORF02236 conserved hypothetical protein
ORF02237 hypothetical protein
ORF02238 hypothetical protein
ORF02239 hypothetical protein
ORF02240 transcriptional regulator, Cro/Cl family
ORF02241 hypothetical protein
ORF02242 transcriptional regulator, Cro/CI family
ORF02243 FtsK/SpollIE family protein
ORF02244 hypothetical protein
ORF02245 hypothetical protein
ORF02246 cell wall surface anchor family protein
ORF02247 transposase, ISL3 family
ORF02250 mercuric resistance operon regulatory protein MerR (merR)
ORF02251 Mn2+/Fe2+ transporter, NRAMP family
ORF02252 membrane protein, putative
ORF02253 ABC transporter, ATP-binding protein
ORF02254 conserved hypothetical protein
ORF02255 streptomycin resistance protein
ORF02257 hypothetical protein
ORF02258 hypothetical protein
ORF02259 conserved hypothetical protein
ORF02260 acetyltransferase, GNAT family
ORF02261 membrane protein, putative
ORF02263 hypothetical protein
ORF02264 transcriptional regulator, Cro/CI family
ORF02265 PAP2 family protein
ORF02266 conserved hypothetical protein FRAMESHIFT
ORF02267 conserved hypothetical protein TIGR00730
ORF02268 protease, putative
ORF02269 rhodanese family protein
ORF02271 hypothetical protein
ORF02274 conserved hypothetical protein
ORF02275 5-methyltetrahydrofolatehomocysteine methyltransferase, putative
ORF02277 conserved hypothetical protein
ORF02279 hypothetical protein
ORF02282 sensor histidine kinase
ORF02283 chromosome assembly-related protein
ORF02287 expressed protein of unknown function
ORF02291 pathogenicity protein, putative
ORF02308 hydrolase, haloacid dehalogenase-like family
ORF02314 conserved hypothetical protein
ORF02317 hypothetical protein
ORF02330 hypothetical protein
ORF02344 site-specific recombinase, phage integrase family
OPEO2245 concound by athetical matein
ORF02345 conserved hypothetical protein
ORF02346 conserved hypothetical protein
ORF02347 hypothetical protein
ORF02349 conserved hypothetical protein
ORF02350 hypothetical protein
ORF02351 transcriptional regulator, Cro/Cl family
ORF02352 conserved domain protein

# Tabl 11: GBS g nes not shared with GAS or pneumoc ccus

ORFxxxxx Ann tation
ORF02354 hypothetical protein
ORF02356 expressed putative secreted protein
ORF02362 sensor histidine kinase
ORF02363 response regulator
ORF02367 membrane protein, putative
ORF02368 conserved hypothetical protein
ORF02379 membrane protein, putative
ORF02395 transcriptional regulator, Cro/CI family
ORF02406 membrane protein, putative
ORF02416 diacylglycerol kinase catalytic domain protein, putative
ORF02418 hypothetical protein
ORF02422 hypothetical protein
ORF02425 conserved hypothetical protein
ORF03001 conserved hypothetical protein
ORF03004 conserved hypothetical protein
ORF03005 cylX protein
ORF03006 Tn916, hypothetical protein
ORF03007 Tn916, hypothetical protein
ORF03008 Tn916, hypothetical protein
ORF03009 Tn916, tetM leader peptide
ORF03010 Tn916, hypothetical protein
ORF03012 prophage LambdaSa2, HNH endonuclease family protein
ORF03013 conserved hypothetical protein
ORF03015 conjugal transfer protein, interruption-N
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# Table 12: GBS ORF's not shared with any publish d genom

UKFXXXXX	Annotati	n
0.550005		

ORF00035 membrane protein, putative
ORF00087 lipoprotein, putative
ORF00088 hypothetical protein
ORF00089 hypothetical protein
ORF00123 hypothetical protein
ORF00138 hypothetical protein
ORF00187 hypothetical protein
ORF00188 hypothetical protein
ORF00192 hypothetical protein
ORF00205 hypothetical protein
ORF00228 lipoprotein, putative
ORF00234 hypothetical protein
ORF00235 hypothetical protein
ORF00238 hypothetical protein
ORF00240 transcriptional regulator, Cro/CI family
ORF00241 hypothetical protein
ORF00242 conserved hypothetical protein
ORF00243 hypothetical protein
ORF00247 hypothetical protein
ORF00249 hypothetical protein
ORF00253 hypothetical protein
ORF00254 hypothetical protein
ORF00255 hypothetical protein
ORF00256 hypothetical protein
ORF00257 hypothetical protein
ORF00258 hypothetical protein
ORF00259 hypothetical protein
ORF00260 hypothetical protein
ORF00272 expressed putative lipoprotein
ORF00273 hypothetical protein
ORF00274 hypothetical protein
ORF00275 hypothetical protein
ORF00276 hypothetical protein
ORF00278 membrane protein, putative
ORF00285 lipoprotein, putative
ORF00292 hypothetical protein
ORF00294 expressed protein of unknown function
ORF00308 conserved hypothetical protein
ORF00332 hypothetical protein
ORF00340 hypothetical protein
ORF00384 hypothetical protein
ORF00402 membrane protein, putative
ORF00408 hypothetical protein
ORF00416 hypothetical protein
ORF00417 hypothetical protein
ORF00448 hypothetical protein
ORF00476 hypothetical protein
ORF00489 DNA-damage-inducible protein J, putative
ORF00490 hypothetical protein
ORF00491 lipoprotein, putativ
ORF00497 conserved domain protein
ORF00510 bacteriocin transport accessory protein, putative
ORF00512 hypothetical protein
ORF00527 hypothetical protein
ORF00556 hypoth tical protein

## Table 12: GBS ORF's not shared with any published g nom

ORFxxxxx Ann tation
ORF00575 hypothetical protein
ORF00599 hypothetical protein
ORF00618 hypothetical protein
ORF00620 hypothetical protein
ORF00623 hypothetical protein
ORF00626 prophage LambdaSa1, transcriptional regulator, Cro/Cl family
ORF00628 hypothetical protein
ORF00630 hypothetical protein
ORF00632 hypothetical protein
ORF00635 hypothetical protein
ORF00636 hypothetical protein
ORF00637 hypothetical protein
ORF00642 conserved hypothetical protein
ORF00644 hypothetical protein
ORF00645 hypothetical protein
ORF00647 hypothetical protein
ORF00649 hypothetical protein
ORF00650 hypothetical protein
ORF00653 conserved hypothetical protein
ORF00657 conserved hypothetical protein, truncation
ORF00661 conserved hypothetical protein
ORF00673 hypothetical protein
ORF00674 hypothetical protein
ORF00675 conserved hypothetical protein
ORF00676 conserved hypothetical protein
ORF00682 hypothetical protein
ORF00685 conserved hypothetical protein
ORF00698 hypothetical protein
ORF00712 hypothetical protein
ORF00718 cell wall surface protein, interruption-N
ORF00723 hypothetical protein
ORF00735 expressed protein of unknown function
ORF00737 conserved hypothetical protein, degenerate
ORF00738 hypothetical protein
ORF00740 hypothetical protein
ORF00741 hypothetical protein
ORF00747 cylD protein (cylD)
ORF00753 cylE protein (cylE)
ORF00756 cylJ protein (cylJ)
ORF00757 cylK protein (cylK)
ORF00758 hypothetical protein
ORF00759 putative secreted protein
ORF00761 hypothetical protein
ORF00796 hydrolase, haloacid dehalogenase-like family
ORF00806 conserved hypothetical protein
ORF00822 ABC transporter, ATP-binding protein
ORF00827 hypothetical protein
ORF00872 cell wall surface anchor family protein
ORF00909 hypothetical protein
ORF00923 hypothetical protein
ORF00924 conserved hypothetical protein
ORF00942 expressed putative secreted protein
ORF00943 hypothetical protein
ORF00944 hypothetical protein
ORF01013 hypothetical protein
Latin a to 10 hypothesiscal protein

#### Tabl 12: GBS ORF's n t shared with any published g nom

ORFxxxxx Ann tation
ORF01014 hypothetical protein
ORF01015 hypothetical protein
ORF01016 hypothetical protein
ORF01018 hypothetical protein
ORF01019 hypothetical protein
ORF01021 hypothetical protein
ORF01035 Tn916, excisionase
ORF01062 hypothetical protein
ORF01096 nisin-resistance protein, putative
ORF01145 lipoprotein, putative
ORF01146 conserved hypothetical protein, FRAMESHIFT
ORF01148 lipoprotein, putative
ORF01149 hypothetical protein
ORF01150 hypothetical protein
ORF01151 hypothetical protein
ORF01152 lipoprotein, putative
ORF01153 hypothetical protein
ORF01158 hypothetical protein
ORF01159 hypothetical protein
ORF01161 expressed conserved domain protein
ORF01162 conserved hypothetical protein
ORF01166 hypothetical protein
ORF01168 conserved hypothetical protein
ORF01169 hypothetical protein
ORF01174 conserved domain protein
ORF01175 hypothetical protein
ORF01186 cell wall surface anchor family protein, putative
ORF01187 hypothetical protein
ORF01204 hypothetical protein
ORF01215 hypothetical protein
ORF01258 hypothetical protein
ORF01262 conserved hypothetical protein, FRAMESHIFT
ORF01263 hypothetical protein
ORF01265 hypothetical protein
ORF01266 hypothetical protein
ORF01304 polysaccharide biosynthesis protein CpsK(V) (cpsK)
ORF01308 polysaccharide biosynthesis protein CpsM(V) (cpsM)
ORF01309 polysaccharide biosynthesis protein cpsH(V) (cpsH)
ORF01349 hypothetical protein
ORF01384 hypothetical protein
ORF01385 hypothetical protein
ORF01386 conserved hypothetical protein
ORF01392 hypothetical protein
ORF01395 conserved hypothetical protein
ORF01409 conserved hypothetical protein
ORF01410 hypothetical protein
ORF01417 hypothetical protein
ORF01418 hypothetical protein
ORF01420 hypothetical protein
ORF01423 conserved hypothetical protein
ORF01424 conserved hypothetical protein
ORF01425 conserved hypothetical protein
ORF01426 conserved hypothetical protein
ORF01427 hypothetical protein
ORF01431 hypothetical protein

#### Table 12: GBS ORF's not shared with any publish d gen me

ORFxxxxx Ann tati n
ORF01432 conserved domain protein
ORF01434 hypothetical protein
ORF01435 calcium-binding protein, putative
ORF01437 abortive infection protein AbiGI (abiGI)
ORF01438 abortive infection protein AbiGI (abiGI)
ORF01441 conserved hypothetical protein, degenerate
ORF01443 hypothetical protein
ORF01445 hypothetical protein
ODE04450 hungath attack markets
ORF01452 hypothetical protein
ORF01463 hypothetical protein
ORF01464 hypothetical protein
ORF01465 hypothetical protein
ORF01486 hypothetical protein
ORF01488 R5 protein
ORF01575 membrane protein, putative
ORF01581 lipoprotein, putative
ORF01601 hypothetical protein
ORF01611 hypothetical protein
ORF01638 conserved hypothetical protein
ORF01645 cell wall surface anchor family protein
ORF01660 membrane protein, putative
ORF01666 hypothetical protein
ORF01667 hypothetical protein
ORF01670 hypothetical protein
ORF01673 hypothetical protein
ORF01674 hypothetical protein
ORF01675 hypothetical protein
ORF01681 hypothetical protein
ORF01682 hypothetical protein
ORF01684 hypothetical protein
ORF01719 hypothetical protein
ORF01733 hypothetical protein
ORF01735 hypothetical protein
ORF01736 hypothetical protein
ORF01737 hypothetical protein
ORF01742 hypothetical protein
ORF01754 hypothetical protein
ORF01761 transposase, IS30 family, putative, truncation
ORF01807 hypothetical protein
ORF01836 hypothetical protein
ORF01838 hypothetical protein
ORF01842 hypothetical protein
ORF01850 hypothetical protein
ORF01892 hypothetical protein
ORF01893 hypothetical protein
ORF01895 hypothetical protein
ORF01896 hypothetical protein
ORF01897 hypothetical protein
ORF01898 hypothetical protein
ORF01899 hypothetical protein
ORF01905 hypothetical protein
ORF01934 hypothetical protein
ORF01943 hypothetical protein
ORF01944 hypothetical protein

## Tabl 12: GBS ORF's not shar d with any published genome

ORFxxxxx Annotation
ORF01953 hypothetical protein
ORF01984 hypothetical protein
ORF01988 hypothetical protein
ORF01989 hypothetical protein
ORF02005 hypothetical protein
ORF02011 hypothetical protein
ORF02016 hypothetical protein
ORF02025 hypothetical protein
ORF02026 hypothetical protein
ORF02045 conserved hypothetical protein
ORF02047 prophage LambdaSa2, holin, putative
ORF02048 conserved hypothetical protein
ORF02049 hypothetical protein
ORF02050 conserved domain protein
ORF02053 conserved hypothetical protein
ORF02057 hypothetical protein
ORF02058 hypothetical protein
ORF02061 hypothetical protein
ORF02062 hypothetical protein
ORF02063 conserved domain protein
ORF02067 conserved hypothetical protein
ORF02069 hypothetical protein
ORF02070 hypothetical protein
ORF02072 conserved hypothetical protein
ORF02073 prophage LambdaSa2, transcriptional regulator, Cro/CI family
ORF02075 hypothetical protein
ORF02077 hypothetical protein
ORF02078 conserved hypothetical protein
ORF02081 hypothetical protein
ORF02085 hypothetical protein
ORF02087 hypothetical protein
ORF02088 conserved hypothetical protein
ORF02091 conserved domain protein
ORF02092 hypothetical protein
ORF02093 hypothetical protein
ORF02094 hypothetical protein
ORF02097 hypothetical protein
ORF02100 hypothetical protein
ORF02102 hypothetical protein
ORF02108 hypothetical protein
ORF02119 hypothetical protein
ORF02124 hypothetical protein
ORF02171 membrane protein, putative
ORF02172 hypothetical protein
ORF02173 membrane protein, putative
ORF02191 hypothetical protein
ORF02196 hypothetical protein
ORF02203 hypothetical protein
ORF02208 hypothetical protein
ORF02212 hypothetical protein
ORF02213 hypothetical protein
ORF02214 transcriptional regulator, Cro/Cl family
ORF02215 expressed protein of unknown function
ORF02217 conserved hypothetical protein
ORF02219 hypothetical protein
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# Tabl 12: GBS ORF's n t shared with any published g nom

#### ORFxxxxx Ann tati n

ORFXXXX AIIII tati II
ORF02221 cell wall anchor protein-related protein
ORF02223 hypothetical protein
ORF02224 hypothetical protein
ORF02225 hypothetical protein
ORF02231 conserved hypothetical protein
ORF02235 hypothetical protein
ORF02236 conserved hypothetical protein
ORF02237 hypothetical protein
ORF02238 hypothetical protein
ORF02239 hypothetical protein
ORF02241 hypothetical protein
ORF02244 hypothetical protein
ORF02245 hypothetical protein
ORF02263 hypothetical protein
ORF02268 protease, putative
ORF02271 hypothetical protein
ORF02279 hypothetical protein
ORF02283 chromosome assembly-related protein
ORF02317 hypothetical protein
ORF02330 hypothetical protein
ORF02344 site-specific recombinase, phage integrase family
ORF02345 conserved hypothetical protein
ORF02347 hypothetical protein
ORF02349 conserved hypothetical protein
ORF02350 hypothetical protein
ORF02351 transcriptional regulator, Cro/CI family
ORF02354 hypothetical protein
ORF02356 expressed putative secreted protein
ORF02395 transcriptional regulator, Cro/CI family
ORF02418 hypothetical protein
ORF02422 hypothetical protein
ORF02425 conserved hypothetical protein
ORF03004 conserved hypothetical protein
ORF03005 cylX protein
ORF03006 Tn916, hypothetical protein
ORF03007 Tn916, hypothetical protein
ORF03008 Tn916, hypothetical protein
ORF03009 Tn916, tetM leader peptide
ORF03010 Tn916, hypothetical protein
ORF03015 conjugal transfer protein, interruption-N

SEQ ID NO. 1301: SAG0466 FROM THE 2603V/R GBS STRAIN

CTCCTGCCCTGCAATGGCAGTTAGACCCATAGGTTTATTTTTATATTTTAATGCCTGCATAAGATGAAGGATATTAATA ATGATTAAATAGAGCATCAATCGCTGCAAATGGTTCATTCCATTCAATTGCATCATAATCCGATATTTTAGTATGAGTTT CTGTTAATAGTTTTTCCGTAGCCGTGTGAACCAATTCTGGACTAAGCTTGGGATCTCCTGCTACTTCTACAATGTGAACA CAAAGGTGAATTTTCCATCAATCTTGGTAATTTTTGAAAAAATGTTtCTTTaGTTTTCTAACGCCTTGATCTCGCATCC CTTCCATTGGTAAGATTACyTCTTCTAAATAGCCACCTTGTTTAGCTGTTAAGGCGCGTTTATGGCTCAAGAATGCCAAT TTATCTAACATTTCTCTTAAAACCATATTTTTGACAGACTCTCTGGGCCCCTTCTAACATTACAGTTTCAGCATAAGA GTCAGGAGAAAACTGAGCAACTGTATATTCTCCGTTACGATTATCTTCTTTAGCATAACGTCTCATAGGTTGAAGAGAAC TACTTTCAATCCCCCCAACAAGAACTTTTTCATTAATACCGGTACTGATTTTTAGATAACCAAAAAACAAGGCAGAACTT GATGAAGCACACTGCATATCAATCGTTTGTACTGGAATATAGGATTCATAATCAGAAAAAAGAGTCATCAAACGACCAAT AAAGGTGTGCTCCTAAAAGTTCTGGACGGTAAGTTTAAATTGCTT

SEQ ID NO. 1302: SAG0466 FROM THE M732 GBS TYPE III STRAIN

GAATCAGAATCTAATATTGATAGTATTATTTGTGGGAACACAGTTGGTACTGGGGGGCAATATTGGTCGTTTGATGACTCT TTTTTCTGATTATGAATCCTATATTCCAGTACAAACGATTGATATGCAGTGTGCTTCATCAAGTTCTGCCTTGTTTTTTG GTTATCTAAAAATCAGTGCCGGTATTAATGAAAAAGTTCTTGTTGGGGGGGATTGAAAGTAGTTCTCTTCAACCTATGAGA CGTTACGCTAAAGAAGATAATCGTAACGGAGAATATACCGTTGCTCAGTTTTCTCCTGACTCTTATGCTGAAACTGTAAT GTTAGAAGGGCACAAAGAGTCTGTCAAAAATATGGTTTTAGAAGAGAAATGTTAGATAAATTGGCATTCTTGAGCCATA AACGCCCCTTAACAGCTAAACAAGGTGGCTATTTAGAAGAGGTAATCTTACCAATGGAAGGGATGCGAGATCAAGGCGTT AGAAAACTAAAAGAAGCATTTTTTCAAAAATTACCAAGATTGATGGAAAATTCACCTTTGCTCACTATTGGAAATGTTTG TTTAATGCACGATGCTGCATTTCTAACGCTTCAGAGTCAGAAAACAGAATTCCGGATTGTTCACATTGTAGAAGTAG CAGGAGATCCCAAGCTTAGTCCAGAATTGGTTCACACGGCTACGGAAAAACTATTAACAGAAACTCATACTAAAATATCG AAAATTCAATATTTTTGGAGGGGCATTAGCTTACGGACACCCTTATGCCTGCTCAGGAATTA

SEQ ID NO. 1303: SAG0466 FROM THE 090 GBS TYPE IA STRAIN TTGTGGGAACACAGTTGGTACTGGGGGCAATATTGGTCGTTTGATGACTCTTTTTTCTGATTATGAATCCTATATTCCAG TACAAACGATTGATATGCAGTGTGCTTCATCAAGTTCTGCCTTGTTTTTTTGGTTATCTAAAAATCAGTGCCGGTATTAAT GAAAAAGTTCTTGTTGGGGGGATTGAAAGTAGTTCTCTTCAACCTATGAGACGTTACGCTAAAGAAGATAATCGTAACGG AGAATATACCGTTGCTCAGTTTTCTCCTGACTCTTAkGCTGAAACTGTAATGLTAGAAGGGGCACAAAGAGTCTGTCAAA AATATGGTTTLAGAAGAGAAATGTTAGATAAATTGGCATTCTTGAGCCATAAACGCGCCCTTAACAGCTAAACAAGGTGGC TATTTAGAAGAGGTAATCTTACCAATGGAAGGGATGCGAGATCAAGGCGTTAGAAAACTAAAAGAAGCATTTTTTCAAAA ATTACCAAGATTGATGGraaattcacctttgctcactattggaaatgtttgtttaatgcacgatgctgcttctwa CGCTTCAGAGTCAGAAAACAGAATTCCGGATTGTTCACATTGTAGAAGTAGCAGGAGATCCCAAGCTTAGTCCAGAATTG GTTCACACGGCTACGGAAAAACTATTAACAGAAACTCATACTAAAATATCGGATTATGATGCAATTGAATGGAATGAACC ATTTGCAGCGATTGATGCTTTATTTAATCATTATTATCCTGAAGAGAGAAAAAATTCAATATTTTTGGAGGGGCATTAG CTTACGGACACCCTTATGCCTGCTCAGG

SEQ ID NO. 1304: SAG0466 FROM THE COH1 GBS TYPE Ia STRAIN ATCGGTATAAAAGGGAAGCAATTTAAAATTACCGTCCAGAACTTTTAGGAGCACCCTCTTAAATCAAATAAAAAAATA GAATCAGAATCTAATATTGATAGTATTATTTGTGGGAACACAGTTGGTACTGGGGGGCAATATTGGTCGTTTGATGACTCT TTTTTCTGATTATGAATCCTATATTCCAGTACAAACGATTGATATGCAGTGTGCTTCATCAAGTTCTGCCTTGTTTTTTG **GGTATCTAAAAA** 

# SEQ ID NO. 1305 : SAG0466 FROM THE CJB GBS NONTYPEABLE STRAIN REVERSE

 ${\tt CATTTCTAACGCTTCAGAGTCAGAAAACAGAATTCCGGATTGTTCACATTGTAGAAGTAGCAGGAGATCCCAAGCTTAGT}$ CCAGAATTGGTTCACACGGCTACGGAAAAACTATTAACAGAAACTCATACTAAAATATCGGATTATGATGCAATTGAATG GAATGAACCATTTGCAGCGATTGATGCTTTATTTAATCATTATTATCCTGAAGAGAGAAAAATTCAATATTTTTGGAG GGGCATTAGCTTACGGACACCCTTAATGCCTGCTCAGGAATTATTAATATCC

SEQ ID NO. 1306: sag0466 FROM THE CJB110 GBS NONTYPEABLE STRAIN  ${\tt ACCAGAATCTAACATTGATAGTATTATTTGTGGGAACACAGTTGGTACTGGGGGGCAATATTGGTCGTTTGATGACTCTTT}$ TTTCTGATTATGAATCCTATATTC

SEQ ID NO. 1309: SAG0466 FROM THE 18RS21 GBS TYPE II STRAIN
TCGGTATAAAAGGGAAGCAATTTAAACATTACCGTCCAGAACTTTTAGGAGCACACCTTTTAAATCAAATAAAAAAATA
GAATCAGAATCTAACATTGATAGTATTATTTGTGGGAACACAGTTGGTACTGGGGGCAATATTGGTCGTTTGATGACTCT
TTTTTCTGATTATGAATCCTATATTCCAGTACAAACGATTGATATGCAGTGTGCTTCATCAAGTTCTGCCTTGTTTTTTG
GTTATCTAAAAATCAGTACCGGTATTAATGAAAAAGTTCTTGTTGGGGGGATTGAAAGTAGTTCTCTTCAACCTATGAGA
CGTTATGCTAAAGAAGATAATCGTAACGGAGAATATACAGTTGCTCAGTTTTCTCCTGACTCTTATGCTGAAACTGTAAT
GTTAGAAGGGGCCCAGAGAGTCTGTCAAAAAATATGGTTTTAGAAAGAGAAATGTTAGATAAATTGGCATTCTTGAGCCATA
AACGCGCCCTTAACAGCTAAACA

SEQ ID NO. 1310: SAG0466 FROM THE H36b GBS TYPE Ib STRAIN.

TTTGGGCTACGAACACCTATCGGTATAAAAGGGAAGCAATTTAAACATTACCGTCCAGAACTTTTAGGAGCACACCTTTT
AAATCAAATAAAAAAATAGAATCAGAATCTAACATTGATAGTATTATTTTGTGGGAACACAGTTGGTACTGGGGGCAATA
TTGGTCGTTTGATGACTCTTTTTTCTGATTATGAATCCTATATTCCAGTACAAACGATTGATATGCAGTGTCTTCATCA
AGTTCTGCCTTGTTTTTTGGTTATCTAAAAATCAGTACCGGTATTAATGAAAAAGTTCTTGTTGGGGGGGATTGAAAGTAG
TTCTCTTCAACCTATGAGACGTTATGCTAAAGAAGATAATCGTAACGGAGAATATACAGTTGCTCAGTTTTCTCCTGACT
CTTATGCTGAAACTGTAATGTTAGAAGGGGCCC

SEQ ID NO. 1312: SAG0466 FROM THE M781 GBS TYPE III STRAIN REVERSER COMPLEMENT CCTTTGCTCACTATTGGAAATGTTTGTTTAATGCACGATGCTGCTGCATTTCTAACGCTTCAGAGTCAGAAACAGAATT CCGGATTGTTCACACTGTAGAAGTAGCAGGAGATCCCAAGCTTAGTCCAGAATTGGTTCACACGGCTACGGAAAAACTAT TAACAGAAACTCATACTAAAATATCGGATTATGATGCAATTGAATGGAATGAACCATTTGCAGCGATTGATGCTTTATTT AATCATTATTATCCTGAAGAGAGAAAAATTCAATATTTTTTGGAGGGGCCATTAGCTTACGGACACCCTTATGCCTGCTC AGGAATTATTAATATCCTTCATCTTTATGCAGGCCATTAAAATATAAAAATAAAACCTATGGGTTCTAACTGC

SEQ ID NO. 1315: SAG0466 FROM THE JM9130013 GBS TYPE VIII STRAIN REVERSE COMPLEMENT

GCTCACTATTGGAAATGTTTGTTTAATGCACGATGCTGCTGCATTTCTAACGCTTCAGAGTCAGAAAACAGAATTCCGGA TTGTTCACATTGTAGAAGTAGCAGGAGATCCCAAGCTTAGTCCAGAATTGGTTCACACGGCTACGGAAAAACTATTAACA GAAACTCATACTAAAATATCGGATTATGATGCAATTGAATGGAATGAACCATTTGCAGCGATTGATGCTCTATTTAATCA TTATTATCCTGAAGAGAGAGAAAAATTCAATATTTTTTGGAGGGGCATTAGCTTACGGACACCCTTATGCCTGCTCAGGAA TTATTAATATCCTTCATCTTATGCAGGCATTAAAATATAAAAATAAAACCTATGGGTCTAACTGCCATTGCAGGGGCAGGA

SEQ ID NO. 1316: SAG0466 FROM THE JM9130013 GBS TYPE VIII STRAIN
TTTGGGCTACGAACACCTATCGGTATAAAAGGGAAGCAATTTAAACATTACCGTCCAGAACTTTTAGGAGCACACCTTTT
AAATCAAATAAAAAAATAGAATCAGAATCTAACATTGATAGTATTATTTGTGGGAACACAGTTGGTACTGGGGGCAATA
TTGGTCGTTTGATGACTCTTTTTTCTGATTATGAATCCTATATTCCAGTACAAACGATTGATATGCAGTGTGCTTCATCA
AGTTCTGCCTTGTTTTTTTGGTTATCTAAAAATCAGTACCGGTATTAATGAAAAAGTTCTTGTTGGGGGGGATTGAAAGTAG
TTCTCTTCAACCTATGAGACGTTATGCTAAAGAAGATAATCGTAACGGAGAATATA

SEQ1301	CTCCTGCCCCTGCAATGGCAGTTAGACCCATAGGTTTATTTTTATATTTTA
SEQ1302	
SEQ1303	
SEQ1304	
SEQ1305	
SEQ1306	
SEQ1307	
SEQ1308	CTTAACAGTTAAACAAGGTGGCTATTTAGAAGAGGTAATCTTACCAATGGAAGGGATGC
SEQ1309	
SEQ1310	
SEQ1311	
SEQ1312	
SEQ1313	
SEQ1314	
SEQ1315	
SEQ1316	
2501210	
SEQ1301	TGCCTGCATAAGATGAAGGATATTAATAATTCCTGAGCAGGCATAAGGGTGTCCGTAAG
SEQ1302	TCGGTATAAA
SE01303	
SEQ1304	ATCGGTATAAA
SE01305	TTTTCAAAAATTACCAAGATTGATGG
SEQ1306	GGTATAA
SEQ1307	CAAGATTGATGG
SEQ1308	AGATCAAGGCGTTAGAAAACTAAAAGAAACATTTTTTCAAAAATTACCAAGATTGATGG
SEQ1309	ТСССТАТАА
SEQ1310	TTTGGGCTACGAACACCTATCGGTATAAA
SEQ1311	G
SEQ1312	
SEQ1313	
SEQ1314	
SEQ1314 SEQ1315	
SEQ1315	TTTGGGCTACGAACACCTATCGGTATAAA
.2FÖ1310	
SEQ1301	TAATGTCCCTCCAAA-AATATTGAATTTTTCTCTCTC-TTCAGGATAATAATGATTAAA
SEQ1302	GGGAAGCAATTTAAACATTACCGTCCAGAACTTTTAGGAGCACACCTCTTAAATCAAAT
SEQ1302 SEQ1303	GOMMONN I INVICTION OF THE CONTROL O
SEQ1303 SEQ1304	GGGAAGCAATTTAAA-ATTACCGTCCAGAACTTTTAGGAGCACACCTCTTAAATCAAAT
SEQ1305	AAATTCACCTTTGCTCACTATTGGAAATGTTTGTTTAATGCACGATGCTGCATTTC
SEQ1306	GGGAAGCAATTTAAACATTACCGTCCAGAACTTTTAGGAGCACACCTCTTAAATCAAAT
	AAATTCACCTTTGCTCACTATTGGAAATGTTTGTTTAATGCACGATGCTGCATTTC
SEQ1307	AAATTCACCTTTGCTCACTATTGGAAATGTTTGTTTAATGCACGATGCTGCATTTC AAATTCACCTTTGCTCACTATTGGAAATGTTTGTTTAATGCACGATGCTGCTGCATTTC
SEQ1308	AAATTCACCTTTGCTCACTATTGGAAATGTTTGTTTAATGCACGATGCTGCTGCATTC
SEQ1309	GGGAAGCAATTTAAACATTACCGTCCAGAACTTTTAGGAGCACACCTTTTAAATCAAAT
SEQ1310	GGGAAGCAATTTAAACATTACCGTCCAGAACTTTTAGGAGCACACCTTTTAAATCAAAT
SEQ1311	AAATTCACCTTTGCTCACTATTGGAAATGTTTGTTTAATGCACGATGCTGCATTTC
SEQ1312	CCTTTGCTCACTATTGGAAATGTTTGTTTAATGCACGATGCTGCATTTC
SEQ1313	GCAATTTAAACATTACCGTCCAGAACTTTTAGGAGCACACCTCTTAAATCAAAT
SEQ1314	CCTTTGCTCACTATTGGAAATGTTTGTTTAATGCACGATGCTGCATTTC
SEQ1315	GCTCACTATTGGAAATGTTTGTTTAATGCACGATGCTGCATTTC

SEQ1316	GGGAAGCAATTTAAACATTACCGTCCAGAACTTTTAGGAGCACACCTTTTAAATCAAAT
-	
SEQ1301	AGAGCATCAATCGCTGCAAATGGTTCATTCC-ATTCAATTGCATCATAATCCGATATTT
SEQ1302	AAAAAAATAGAATCAGAATCTAATATTGATAGTATTATTTGTGGGAACA-CAGT
SEQ1303	TTGTGGGAACA-CAGT
SEQ1304	AAAAAATAGAATCAGAATCTAATATTGATAGTATTATTTGTGGGAACA-CAGT
SEQ1305	AACGCTTCAGAGTCAGAAAACAGAATTCCGGATTGTTCACATTGTAGAAGTAGCAGG
SEQ1306	AAAAAATATAACCAGAATCTAACATTGATAGTATTATTTGTGGGAACA-CAGT
SEQ1307	AACGCTTCAGAGTCAGAAAACAGAATTCCGGATTGTTCACATTGTAGAAGTAGCAGG
SEQ1308	AACGCTTCAGAGTCAGAAAACAGAATTCCGGATTGTTCACATTGTAGAAGTAGCAGG
SEQ1309	AAAAAATAGAATCAGAATCTAACATTGATAGTATTATTTGTGGGAACA-CAGT
SEQ1310	AAAAAAATAGAATCAGAATCTAACATTGATAGTATTATTTGTGGGAACA-CAGT
SEQ1311	AACGCTTCAGAGTCAGAAAACAGAATTCCGGATTGTTCACATTGTAGAAGTAGCAGG
SEQ1312	AACGCTTCAGAGTCAGAAAACAGAATTCCGGATTGTTCACATTGTAGAAGTAGCAGG
SEQ1313	Aaaaaatagaatcagaatctaatattgatagtattatttgtgggaaca-cagt
SEQ1314	AACGCTTCAGAGTCAGAAAACAGAATTCCGGATTGTTCACATTGTAGAAGTAGCAGG
SEQ1315	AACGCTTCAGAGTCAGAAAACAGAATTCCGGATTGTTCACATTGTAGAAGTAGCAGG
SEQ1316	AAAAAATAGAATCAGAATCTAACATTGATAGTATTATTTGTGGGAACA-CAGT
SEQ1301	AGTATGAGTTTCTGTTAATAGTTTTTCCGTAGCCGTGTGAACCAATTCTGGACTAAGCT
SEQ1302	GGTACTGGGGGCAATATTGG-TCGTTTGATGACTCTTTTTTCTGATTATGAATCCTA
SEQ1303	GGTACTGGGGGCAATATTGG-TCGTTTGATGACTCTTTTTTCTGATTATGAATCCTA
SEQ1304	GGTACTGGGGGCAATATTGG-TCGTTTGATGACTCTTTTTTCTGATTATGAATCCTA
SEQ1305	GATCCCAAGCTTAGTCCAGAATTGGTTCACACGGCTACGGAAAAACTATTAACAGAAAC
SEQ1306	GGTACTGGGGGCAATATTGG-TCGTTTGATGACTCTTTTTTCTGATTATGAATCCTA
SEQ1307	GATCCCAAGCTTAGTCCAGAATTGGTTCACACGGCTACGGAAAAACTATTAACAGAAAC
SEQ1308	GATCCCAAGCTTAGTCCAGAATTGGTTCACACGGCTACGGAAAAACTATTAACAGAAAC
SEQ1309	GGTACTGGGGGCAATATTGG-TCGTTTGATGACTCTTTTTTCTGATTATGAATCCTA
SEQ1310	GGTACTGGGGGCAATATTGG-TCGTTTGATGACTCTTTTTTCTGATTATGAATCCTA
SEQ1311	GATCCCAAGCTTAGTCCAGAATTGGTTCACACGGCTACGGAAAAACTATTAACAGAAAC
SEQ1312	GATCCCAAGCTTAGTCCAGAATTGGTTCACACGGCTACGGAAAAACTATTAACAGAAAC
SEQ1313	GGTACTGGGGGCAATATTGG-TCGTTTGATGACTCTTTTTTCTGATTATGAATCCTA
SEQ1314	GATCCCAAGCTTAGTCCAGAATTGGTTCACACGGCTACGGAAAAACTATTAACAGAAAC
SEQ1315	GATCCCAAGCTTAGTCCAGAATTGGTTCACACGGCTACGGAAAAACTATTAACAGAAAC
SEQ1316	GGTACTGGGGGCAATATTGG-TCGTTTGATGACTCTTTTTTCTGATTATGAATCCTA
<b></b>	
SEQ1301	GGGATCTCCTGCTACTTCTACAATGTGAACAATCCGGA-ATTCTGTTTTCTGACTCTGA
SEQ1302	TATTCCAGTACAAACGATTGATATGCAGTGTGCTTCATCAAGTTCTGCCTTGTTTTT
:SEQ1303	TATTCCAGTACAAACGATTGATATGCAGTGTGCTTCATCAAGTTCTGCCTTGTTTTT
SEQ1304	TATTCCAGTACAAACGATTGATATGCAGTGTGCTTCATCAAGTTCTGCCTTGTTTTT
SEQ1305	CATACTAAAATATCGGATTATGATGCAATTGAATGGAATGAACCATTTGCAGCGATTGA
SEQ1306	TATTC
SEQ1307	CATACTAAAATATCGGATTATGATGCAATTGAATGGAATGAACCATTTGCAGCGATTGA
SEQ1308	CATACTAAAATATCGGATTATGATGCAATTGAATGGAATGAACCATTTGCAGCGATTGA
SEQ1309	TATTCCAGTACAAACGATTGATATGCAGTGTGCTTCATCAAGTTCTGCCTTGTTTTT
SEQ1310	TATTCCAGTACAAACGATTGATATGCAGTGTGCTTCATCAAGTTCTGCCTTGTTTTT
SEQ1311	CATACTAAAATATCGGATTATGATGCAATTGAATGGAATGAACCATTTGCAGCGATTGA
SEQ1312	CATACTAAAATATCGGATTATGATGCAATTGAATGGAATGAACCATTTGCAGCGATTGA
SEQ1313	TATTCCAGTACAAACGATTGATATGCAGTGTGCTTCATCAAGTTCTGCCTTGTTTTT
SEQ1314	CATACTAAAATATCGGATTATGATGCAATTGAATGGAATGAACCATTTGCAGCGATTGA
SEQ1315	CATACTAAAATATCGGATTATGATGCAATTGAATGGAATGAACCATTTGCAGCGATTGA
SEQ1316	TATTCCAGTACAAACGATTGATATGCAGTGTGCTTCATCAAGTTCTGCCTTGTTTTT
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SEQ1301	GCGTTAGAAATGCAGCAGCATCGTGCATTAAACAAACATTTCCAATAGTGAGCAAAG
SEQ1302	GGT-TATCTAAAAATCAGTG-CCGGTATTAATGAAAAAGTTCTTGTTGGGGGGGATTGAA
SEQ1303	GGT-TATCTAAAAATCAGTG-CCGGTATTAATGAAAAAGTTCTTGTTGGGGGGGATTGAA
SEQ1304	GGG-TATCTAAAAA
SEQ1305	GCTTTATTTAATCATTATTATCCTGAAGAGAGAAAAATTCAATATTTTTTGGAGGGGC
SEQ1306	
SEQ1307	GCTTTATTTAATCATTATTATCCTGAAGAGAGAGAAAAATTCAATATTTTTGGAGGGGC
SEQ1308	GCTCTATTTAATCATTATTATCCTGAAGAGAGAAAAATTCAATATTTTTGGAGGGAC
SEQ1309	GGT-TATCTAAAAATCAGTA-CCGGTATTAATGAAAAAGTTCTTGTTGGGGGGATTGAA
SEQ1310	GGT-TATCTAAAAATCAGTA-CCGGTATTAATGAAAAAGTTCTTGTTGGGGGGATTGAA
SEQ1311	GCTCTATTTAATCATTATTATCCTGAAGAGAGAGAAAAATTCAATATTTTTGGAGGGAC
SEQ1312	GCTTTATTTAATCATTATTATCCTGAAGAGAGAGAAAAATTCAATATTTTTGGAGGGGC
SEQ1313	GGT-TATCTAAAAATCAGTG-CCGGTATTAATGAAAAAGTTCTTGTTGGGGGGATTGAA
SEQ1314	GCTCTATTTAATCATTATTATCCTGAAGAGAGAGAAAAATTCAATATTTTTTGGAGGGAC
SEQ1315	GCTCTATTTAATCATTATTATCCTGAAGAGAGAGAAAAATTCAATATTTTTGGAGGGGC
SEQ1316	GGT-TATCTAAAAATCAGTA-CCGGTATTAATGAAAAAGTTCTTGTTGGGGGGATTGAA
SEQ1301	TGAATTTTCCATCAATCTTGGTAATTTTTGAAAAAATGTTTCTTTTAGTTTTCTAAC
SEQ1302	GTAGTTCTCTTCAACCTATGAGACGTTACGCTAAAGAAGATAATCGTAACGGAGAATAT
SEQ1303	GTAGTTCTCTTCAACCTATGAGACGTTACGCTAAAGAAGATAATCGTAACGGAGAATAT
SEQ1304	THE TOTAL TOTAL TOTAL THE TOTAL THE TOTAL TOTAL TOTAL TOTAL THE TO
SEQ1305	TTAGCTTACGGACACCCTTAATGCCTGCTCAGGAATTATTAATATCC
SEQ1306	**************************************
SEQ1307	TTAGCTTACGGACACCCTTATGCCTGCTCAGGAATTATTAATATCCTTCATCTTAT
SEQ1308	TTAGCTTACGGACACCCTTATGCCTGCTCAGGAATTATTAATATCCTTCATCTTAT
SEQ1309	GTAGTTCTCTTCAACCTATGAGACGTTATGCTAAAGAAGATAATCGTAACGGAGAATAT
SEQ1310	GTAGTTCTCTTCAACCTATGAGACGTTATGCTAAAGAAGATAATCGTAACGGAGAATAT
SEQ1311	TTAGCTTACGGACACCCTTATGCCTGCTCAGGAATTATTAATATCCTTCATCTTAT
SEQ1312	TTAGCTTACGGACACCCTTATGCCTGCTCAGGAATTATTAATATCCTTCATCTTAT
SEQ1313	GTAGTTCTCTTCAACCTATGAGACGTTACGCTAAAGAAGATAATCGTAACGGAGAATAT
SEQ1314	TTAGCTTACGGACACCCTTATGCCTGCTCAGGAATTATTAATATCCTTCATCTTAT
SEQ1315	TTAGCTTACGGACACCCTTATGCCTGCTCAGGAATTATTAATATCCTTCATCTTAT
SEQ1316	GTAGTTCTCTTCAACCTATGAGACGTTATGCTAAAGAAGATAATCGTAACGGAGAATAT
SEQ1301	CCTTGATCTCGCATCCCTTCCATTGGTAAGATTACYTCTTCTAAATAGCCACCTTGTTT
SEQ1302	CCGTTGCTCAGTTTTCTCCTGACTCTTATGCTGAAACTGTAATGTTAGAAGGGGCAC
SEQ1303	CCGTTGCTCAGTTTTCTCCTGACTCTTAKGCTGAAACTGTAATGTTAGAAGGGGCAC
SEQ1304	
SEQ1305	
SEQ1306	
SEQ1307	CAGGCATTAAAATAAAAATAAACCTATGGGC-CTAACTGCCATTGCAGGGGCA
SEQ1308	CAGGCATTAAAATAAAAATAAACCTATGGGT-CTAACTGCCATTGCAGGGGCAG
SEQ1309	CAGTTGCTCAGTTTTCTCCTGACTCTTATGCTGAAACTGTAATGTTAGAAGGGGCCC
SEQ1310	CAGTTGCTCAGTTTTCTCCTGACTCTTATGCTGAAACTGTAATGTTAGAAGGGGCCC
SEQ1311	CAGGCATTAAAATATAAAATAAACCTATGGGT-CTAACTGCCATTGCAGGGGCAGGA-
SEQ1312	CAGGCATTAAAATATAAAAATAAACCTATGGGTTCTAACTGC
SEQ1313	CCGTTGCTCAGTTTTCTCCTGACTCTTATGCTGAAACTGTAATGTTAGA
SEQ1314	CAGGCATTAAAATATAAAATAAACCTATGGGT-CTAACTGCCATTGCAGGGGC
SEQ1315	CAGGCATTAAAATATAAAAATAAACCTATGGGT-CTAACTGCCATTGCAGGGGCAGGA-
SEQ1316	TABCMARATVSTNCSRATNGTSAGTHAS

GCTGTTAAGGCGCGTTTATGGCTCAAGAATGCCAATTTATCTAACATTTCTCTTAAA SEQ1301 AAGAGTCTGTCAAAAATATGGTTTTAGAAGAGAAATGTTAGATAAATTGGCATTCTTGA SEQ1302 SEQ1303 AAGAGTCTGTCAAAAATATGGTTTTAGAAGAGAAATGTTAGATAAATTGGCATTCTTGA SEQ1304 SEQ1305 SEQ1306 SEQ1307 SEQ1308 **SEQ1309** GAGAGTCTGTCAAAAATATGGTTTTAGAAGAGAAATGTTAGATAAATTGGCATTCTTGA \_\_\_\_\_\_ SEQ1310 SEQ1311 SEQ1312 SEQ1313 \_\_\_\_\_\_ SEQ1314 \_\_\_\_\_\_ SEQ1315 SEQ1316 SEQ1301 CCATATTTTTGACAGACTCTCTGGGCCCCTT--CTAACATTACAGTTTCAGCATAAGAG SEQ1302 CCATAAACGCGCCTTAACAGCTAAACAAGGTGGCTATTTAGAAGAGGTAATCTTACCAA SE01303 CCATAAACGCGCCTTAACAGCTAAACAAGGTGGCTATTTAGAAGAGGTAATCTTACCAA **SEQ1304** \_\_\_\_\_\_ **SEQ1305 SEQ1306 SEQ1307** SEQ1308 SEQ1309 CCATAAACGCGCCTTAACAGCTAAACA-----SEQ1310 SEQ1311 SEQ1312 SEQ1313 SEQ1314 SE01315 SEQ1316 SEQ1301 CAGGAGAAAACTGAGCAACTGTATATTCTCCGTTACGATTATCTTCTTTAGCATAACGT SEQ1302 GGAAGGGATGCGAGATCAAGGCGTTAGAAAACTAAAAGAAGCATTTTTTCAAAAATTAC **SEQ1303** GGAAGGGATGCGAGATCAAGGCGTTAGAAAACTAAAAGAAGCATTTTTTCAAAAATTAC **SEQ1304** SEQ1305 SEQ1306 **SEQ1307** SEQ1308 SEQ1309 SEQ1310 SEQ1311 **SEQ1312** SEQ1313 SEQ1314 SEQ1315 SEQ1316

SEQ1301	TCATAGGTTGAAGAACTACTTTCAATCCCCCCAACAAGAACTTTTTCATTAATACCG
SEQ1302	AAGATTGATGGAAAATTCACCTTTGCTCACTATTGGAAATGTTTGTT
SEQ1303	AAGATTGATGGRAAATTCACCTTTGCTCACTATTGGAAATGTTTGTTTAATGCACGATG
SEQ1304	
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SEQ1301	TACTGATTTTTAGATAACCAAAAAACAAGGCAGAACTTGATGAAGCACACTGCATAT
· SEQ1302	TGCTGCATTTCTAACGCTTCAGAGTCAGAAAACAGAATTCCGGATTGTTCACATTGTAG
SEQ1302	TGCTGCATTTCTWACGCTTCAGAGTCAGAAAACAGAATTCCGGATTGTTCACATTGTAG
_	IGCIGCATTICIWACGCTICAGAGICAGAAAACAGAATTCCGGATTGTTCACATTGTAG
SEQ1304	
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SEQ1314	
SEQ1315	
SEQ1316	
SEQ1301	AATCGTTTGTACTGGAATATAGGATTCATAATCAGAAAAAAGAGTCATCAAACGACCAA
SEQ1302	AGTAGCAGGAGATCCCAAGCTTAGTCCAGAATTGGTTCACACGGCTACGGAAAAACTAT
SEQ1303	AGTAGCAGGAGATCCCAAGCTTAGTCCAGAATTGGTTCACACGGCTACGGAAAAACTAT
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SEQ1301	ATTGCCCCCAGTACCAACTGTGTTCCCACAAATAATACTATCAATGTTAGATTCTGATT
SEQ1302	AACAGAAACTCATACTAAAATATCGGATTATGATGCAATTGAATGGAATGAACCATTTG
SEQ1303	AACAGAAACTCATACTAAAATATCGGATTATGATGCAATTGAATGGAATGAACCATTTG
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SEQ1310	#
SEQ1311	
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SEQ1316	
SEQ1301	TATTTTTTTTTTTTGATTTAAAAGGTGTGCTCCTAAAAGTTCTGGACGGTAAGTTTAAA
SEQ1302	AGCGATTGATGCTTTATTTAATCATTATTATCCTGAAGAGAGAG
SEQ1303	AGCGATTGATGCTTTATTTAATCATTATTATCCTGAAGAGAGAG
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SEQ1311	
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SEQ1316	
SEQ1301	TGCTT
SEQ1302	TGGAGGGCATTAGCTTACGGACACCCTTATGCCTGCTCAGGAATTA
SEQ1303 ·	TGGAGGGGCATTAGCTTACGGACACCCTTATGCCTGCTCAGG
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SEQ1315	
SEQ1316	



#### Table 14: C mparative Sequences relating to SAG0471 (glucokinase)

SEQ ID NO. 1402: SAG0471 FROM THE 090 GBS TYPE IA STRAIN
CGTTTCTGATATCGTTGAATCTCTCAAACATCGTTTGAGCCTCTATGGATTAACAAAAGATGACTTTCTCGGTATCGGTATGGGTTCTC
CAGGAGCTGTTGATAGAACTAGTAAAACAGTAACAGGTGCTTTTAATCTAAATTGGGCTGATACTCAAGAAGTAGGTTCGGTTATTGAA
AAAGAAGTTGGAATTCCATTTTTTATTGATAACGATGCTAATGTTGCAGCACTTGGTGAACCCTGGGTAGGTGCTGGTGCCAATAATCC
CGATGTTGTTTTCGTAACCCTCGGAACAGGAGTAGGTGGAGGTGTTATCGCAGATTGGTAACACTCCATGGTGTTTCCAGGAGCAGGTG
GAGAAATTGGGCATATGATTGTTGATCCAGAAATGGATTTACGTGCACATGTGGTAACAAAGGCTGTCTTGAGACAGTTGCATCAGCG
ACAGGTTGTTAGAGTAGCACGTCAACTCGCAGAACAATATGAAGGTTCGTCTGTCGTCATTAAAGCACGTGATACCTTGACTGT
TACAAGTAAAGATATTTTTATAGCAGCAGAAGATTGGGGTTAATTTTGCTAATTCTGTTGTTGAACGTGTAATTACCTTAGCTTGCCC
CAGCAGCTAATATTTCAAATATTTTAAACCCTGATTCTGTGGTTATTTGGTGGCGGTGTCTCAGCAGCAGGTGAATTTTACGTAGTCGC

SEQ ID NO. 1403: SAG0471 FROM THE COH1 GBS TYPE Ia STRAIN
ACAAGAAAAATGGGCAATTGAGACCAATACTTTAGAAAACGGAAGACATATCGTTTCTGATATCGTTGAATCTCTCAAACATCGTTTGA
GCCTCTATGGATTAACAAAAAGATGACTTCTCGGTATCGGTATGGGTTCTCCAGGAGCTGTTGATAGAACTAGTAAAACAGTAACAGGT
GCTTTTAATCTAAATTGGGCTGATACTCAAGA

**GTTGAGAAATACTTTGTCACATTTG** 

SEQ ID NO. 1405: SAG0471 FROM THE CJB110 GBS NONTYPEABLE STRAIN
CACCAGCTAATATTTCAAATATTTTAAACCCTGATTCTGTGGTTATTGGTGGCGGTGTCTCAGCAGCAGGTGAATTTTTACGTAGTCGC
GTTGAGAAATACTTTGTCACATTTGCTTTCCCACAAGTTAAAAAGTCAACTA

SEQ ID NO. 1406: SAG0471 FROM THE 2603V/R GBS TYPE V STRAIN
GGGCAATTGAGACCAATACTTTAGAAAACGGAAGACATATCGTTTCTGATATCGTTGAATCTCTCAAACATCGTTTGAGCCTCTATGGA
TTAACAAAAGATGACTTTCTCGGTATCGGTATCGGTTCTCCAGGAGCTG

SEQ ID NO. 1407: SAG0471 FROM THE H36b GBS TYPE Ib STRAIN
GGCAATTGAGACCAATACTTTAGAAAACGGAAGACATATCGTTTCTGATATCGTTGAATCTCTCAAACATCGTTTGAGCCTCTATGGAT
TAACAAAAGATGACTTTCTCGGTATCGGTATGGGTTCTCCAGGAGCTGTTGATAGAACTAGTAAAACAGTAACAGTGCTTTTAATCTA
AATTGGGCTGATACTCAAGAAGTAGGTTCAGTTATTGAAAAAGAAGTTGGAATTCCATTTTTTATTGATAACGATGCTAATGTTGCAGC
ACTTGGTGAACGCTGGGTAGGTGCTGGTGCCAATAATCCCGACGTTGTTTTCGTAACC

SEQ ID NO. 1408: SAG0471 FROM THE H36 GBS TYPE Ib STRAIN REVERSE COMPLEMENT GAGACAGTTGCATCAGCGACAGGTGTTGTTAGAGTAGCAGCGATTGACACTCGCAGAACAATATGAGGGTTCGTCTGCCATTAAAGCAGCGATTGACAACGGTGATACTGTTACAAGTAAAGATATTTTTATAGCAGCAGAAGATGGGGATAAATTTTGCTAATTCTGTTGTTGAACGTGTATCACGTTACCTTGGACTGGCAGCAGCAGATAATTTTTAAACCCTGATTCTGTGGTTATTGGTGGCGGTGTCTCAGCAGCAGGTGAATTTTTTACGTAGTCGCGTTGAGAAATACTTTGTCACATTTGCTTTCCCACA

SEQ ID NO. 1409: SAG0471 FROM THE M732 GBS TYPE III STRAIN
ACAAGAAAAATGGCCAATTGAGACCATACTTAGAAAACGGAAGACATATCGTTTCTGATATCGTTGAATCTCTCAAACATCGTTTGAGC
CTCTATGGATTAACAAAAGATGACTTTCTCGGTATCGGTATGGGTTCTCCAGGAGCTGTTGATAGAACTAGTAAAACAGTAACAGGTGC
TTTTAATCTAAATTGGGCTGATACTCAAGAAGTAGGTTCGGTTATTGAAAAAGAAGTTGGAATTCCATTTTTTATTGATAACGATGCTA
ATGTTGCAGCACTTGGTGAACGCTGGGTAGGTGCCAATAATCCCGATGTTGTTTTCGTAACCCTCGGAACAGGAGTAGGTGGA
GGTGTTATCGCAGATGGTAACCTCCATCCATGGTGTTTCCAAGAGCAGGTGGAGAAATTGGGCATATGATT

SEQ ID NO. 1410: SAG0471 FROM THE M732 GBS TYPE III STRAIN REVERSE COMPLEMENT CAGCAGCAGGTGAATTTTTACGTAGTCGCGTTGAGAAATACTTTGTCACATTTGCTTTCCCACAAGTTAAAAAGTCAACTAAAATTAAGATTGCTGAACTAGGTAATGAT

SEQ ID NO. 1411: SAG0471 FROM THE M781 GBS TYPE III STRAIN
AGAAGTACAAGAAAATGGGCAATTGAGACCATACTTAGAAAACGGAAGACATATCGTTTCTGATATCGTTGAATCTCTCAAACATCGTT
TGAGCCTCTATGGATTAACAAAAGATGACTTTCTCGGTATCGGTATGGGTTCTCCAGGAGCTGTTGATAGAACTAGTAAAACAGTAACA
GGTGCTTTTAATCTAAATTGGGCTGATACTCAAGAAGTAGGTTCGGTTATTGAAAAAGAAGTTGGAATTCCATTTTTTATTGATAACGA
TGCTAATGTTGCAGCACTTGGTGAACGCTGGGTAGGTGCTGGTGCCAATAATCCCGATGTTGTTTCGTAACCCTCGGAACAGGAGTA



#### Table 14: Comparative Sequences relating to SAG0471 (glucokinase)

SEQ ID NO. 1412: SAG0471 FROM THE M781 GBS TYPE III STRAIN REVERSE COMPLEMENT
GATACTGTTACAAGTAAAGATATTTTTATAGCAGCAGAAGATGGGGATAAATTTGCTAATTCTGTTGTTGAACGTGTATCACGTTACCT
TGGACTGGCAGCAGCTAATATTTCAAATATTTTAAACCCTGATTCTGTGGTTATTGGTGGCGGTGTCTCAGCAGCAGGTGAATTTTTAC
GTAGTCGCGTTGAGAAATACTTTGTCACATTTGCTTTCCCACAAGTTAAAAA

SEQ ID NO. 1413: SAG0471 FROM THE 090 GBS TYPE IA STRAIN
AAATTTGGTATCTTGACGCTTGACGGGAGAAGTACAAGAAAAATGGGCATTGAGACCATACTTAGAAAACGGAAGACATATCGTTTCTGA
TATCGTTGAAACTTCTCAAACACGGTGCTTTTGAGCCTCTATGGATTAACAAAAGATGACTTTCTCGGTATCGGTATCGGTTCTCCAGGAGCTG
TTGATAGAACTAGTAAAACAGTAACAGGTGCTTTTAATCTAAATTGGGCTGATACTCAAGAAGTAGGTTCAGTTATTGAAAAAGAAGTT
GGGAATTCCATTTTTTATTGATAACGATGCTAATGTTGCAGCACTTGGTGAACGCTGGGTAGGTGCTGGTGCCAATAATCCCGACGTTGT
TTTCGTAACCCTCGGAACAGGAGTAGGTGGAGG

SEQ ID NO. 1414: SAG0471 FROM THE 090 GBS TYPE IA STRAIN REVERSE COMPLEMENT GTGATACTGTTACAAGTAAAGATATTTTATAGCAGCAGAAGATGGGGATAAATTTGCTAATTCTGTTGTTGAACGTGTACCCTTACCCTTACCCTTGGACTGGCAGCAGCAAGATATTTTCAAATATTTTAAACCCTGATTCTGTGGTTATTGGTGGCGGTGTCTCAGCAGCAGGTGAATTTTTAAACCCTGATTCTGTGGTTAATAGGTGACTAAAATTAAGATTGACGTGAACTTAAAAAATTAAGATTG

SEQ ID NO. 1415: SAG0471 FROM THE JM9130013 GBS TYPE VIII STRAIN REVERSE COMPLEMENT GTTATCGCAGATGGTAACCTCATCGTGTGTTGCAGGAGCAGGTGGAGAAATTGGGCATATGATTGTTGATCCAGAAAATGGATTTAC GTGCACATGTGGTAACAAAGGCTGCCTTGAGACAGTTGCATCAGCGACAGGTGTTGTTAGAGTAGCACGGTCAACTCGCAGAACAATATG AGGGTTCGTCTGCCATTAAAGCAGCGATTGACCACGGTGATACTGTTACAAGTAAAGATATTTTTATAGCAGCAGAAGATGGGGATAAA TTTGCTAATTCTTGTTGAACGTGTATCACGTTACCTTGGACTGCCAGCAGCAGCTAATATTTTCAAATATTTTAAACCCTGATTCTGTGGTTATTTGGTGGCGGTGTCTCCAGCAGGTGAATTTTTACGTAGTCGCGTTGAGAAATACTTTGTCACATTTGCTTTCCCACAAGTTAAAA AGTCAACTAA

SEQ ID NO. 1416: SAG0471 FROM THE JM9130013 GBS TYPE VIII STRAIN REVERSE COMPLEMENT TGGTATCTTGACGCTTGAGGGAGAGTACAAGAAAAATGGGCAATTGAGACCATACTTAGAAAACGGAAGACATATCGTTTCTGATATC GTTGAATCTCTCAAACATCGTTTGAGCCTCTATGGATTAACAAAAGATGACTTTCTCGGTATCGGTATGGGTTCTCCAGGAGCTGTTGA TAGAACTAGTAAAACAGTCACAGGTGCTTTTAATCTAAATTGGGCTGATACTCAAGAAGTAGGTTCAGTTATTGAAAAAGAAGATGGAA TTCCATTTTTTATTG

SEQ ID NO. 1417: SAG0471 FROM THE 2603V/R TYPE V GBS STRAIN REVERSE COMPLEMENT AGCAGCTAATATTTCAAATATTTTAAACCCTGATTCTGTGGTTATTGGTGGCGGTGTCTCAGCAGCAGGTGAATTTTTACGTAGTCGCG TTGAGAAATACTTTGTCACATTTGTTTTCCCACAAGGT

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SEQ1415	TTATCGCAGATGGTAACCTCATCCATGGTGTTGCAGGAGCAGGTGGAGAAATTGGGCAT
SEQ1416	
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SEQ1402 SEQ1403 SEQ1404 SEQ1405 SEQ1406 SEQ1407 SEQ1408 SEQ1409 SEQ1410 SEQ1411 SEQ1411 SEQ1412 SEQ1412 SEQ1413 SEQ1414 SEQ1415	TGATTGTTGATCCAGAAAATGGATTTACGTGCACATGTGGTAACAAAGGCTGCCTTGAG
SEQ1402 SEQ1403 SEQ1404 SEQ1405 SEQ1406 SEQ1407 SEQ1408 SEQ1409 SEQ1410 SEQ1411 SEQ1411 SEQ1412 SEQ1412 SEQ1414 SEQ1415 SEQ1416	
SEQ1402 SEQ1403 SEQ1404 SEQ1405 SEQ1406 SEQ1407 SEQ1408 SEQ1409 SEQ1410 SEQ1411 SEQ1411 SEQ1412 SEQ1412 SEQ1413 SEQ1414 SEQ1415	

## Tabl 14: Comparative Sequences relating t SAG0471 (glucokinase)

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SEQ1401_	
SEQ1402	
SEQ1403	
SEQ1404	*
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SEQ1406	~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~
SEQ1407	
SEQ1408	CAGTTGCATCAGCGACAGGTGTTGTTAGAGTAGCACGTCAACTCGCAGAACAATATGAG
SEQ1409	The state of the s
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SEQ1411	
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SEQ1413	
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SEQ1415	CAGTTGCATCAGCGACAGGTGTTGTTAGAGTAGCACGTCAACTCGCAGAACAATATGAG
SEQ1416	
SEQ1417	
SEQ1401	
SEQ1402	
SEQ1403	
SEQ1404	
SEQ1405	
SEQ1406	
SEQ1407	
șe <b>Q14</b> 08	GTTCGTCTGCCATTAAAGCAGCGATTGACAACGGTGATACTGTTACAAGTAAAGATATT
SEQ1409	
SEQ1410	
SEQ1411	
SEQ1412	GATACTGTTACAAGTAAAGATATT
SEQ1413	
SEQ1414	GTGATACTGTTACAAGTAAAGATATT
SEQ1415	GTTCGTCTGCCATTAAAGCAGCGATTGACCACGGTGATACTGTTACAAGTAAAGATATT
SEQ1416	CITCOICIGCAITAAAGCAGCGAITGACCACGGIGATACIGITACAAGTAAAGATATT
SEQ1417	
SPATATI	
0001401	
SEQ1401_	TTAAATTTGGTATCTTGACGCTTGAGGGAGAAGTACAA
SEQ1402	
SEQ1403	ACAA
SEQ1404	TTGGTATCTTGACGCTTGAGG-AGAAGTACAA
SEQ1405	
SEQ1406	
SEQ1407	
SE01408	TTATAGCAGCAGAAGATGGGGATAAATTTGCTAATTCTGTTGTTGAACGTGTATCACGT
SEQ1409	ACAA
SEQ1410	ACAA
SEQ1411	AGAAGTACAA
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SEQ1412	TTATAGCAGCAGAAGATGGGGATAAATTTGCTAATTCTGTTGTTGAACGTGTATCACGT
SEQ1413	AAATTTGGTATCTTGACGCTTGAGGGAGAAGTACAA
SEQ1414	TTATAGCAGCAGAAGATGGGGATAAATTTGCTAATTCTGTTGTTGAACGTGTATCACGT
SEQ1415	TTATAGCAGCAGAAGATGGGGATAAATTTGCTAATTCTGTTGTTGAACGTGTATCACGT
SEQ1416	TGGTATCTTGACGCTTGAGGGAGAAGTACAA
SEQ1417	

## Table 14: C mparative Sequences relating t SAG0471 (gluc kinase)

SEQ1401_	AAAAATGGGCAATTGAGACCAATACTTTAGAAAACGGAAGACATATCGTTTCTGATATC
SEQ1402	CGTTTCTGATATC
SEQ1403	AAAAATGGGCAATTGAGACCAATACTTTAGAAAACGGAAGACATATCGTTTCTGATATC
SEQ1404	AAAAATGGGCAATTGAGACCAATACTTTAGAAAACGGAAGACATATCGTTTCTGATATC
SEQ1405	CACCAGCTAATATTTCAAATATTTTAAACCCTGATTCTGTGGTTATT
SEQ1406	GGGCAATTGAGACCAATACTTTAGAAAACGGAAGACATATCGTTTCTGATATC
SEQ1407	GGCAATTGAGACCAATACTTTAGAAAACGGAAGACATATCGTTTCTGATATC
SEQ1408	ACCTTGGACTGGCAGCAATATTTCAAATATTTTAAACCCTGATTCTGTGGTTATT
SEQ1409	AAAAATGGGCAATTGAGACCA-TACTT-AGAAAACGGAAGACATATCGTTTCTGATATC
SEQ1410	
SEQ1411	AAAA-TGGGCAATTGAGACCA-TACTT-AGAAAACGGAAGACATATCGTTTCTGATATC
SEQ1412	ACCTTGGACTGGCAGCAATATTTCAAATATTTTAAACCCTGATTCTGTGGTTATT
SEQ1413	AAAAATGGGCA-TTGAGACCA-TACTT-AGAAAACGGAAGACATATCGTTTCTGATATC
SEQ1414	ACCTTGGACTGGCAGCAATAATTTCAAATATTTAAACCCTGATTCTGTGGTTATT
SEQ1415	ACCTTGGACTGGCAGCTAATATTTCAAATATTTTAAACCCTGATTCTGTGGTTATT
SEQ1416	AAAAATGGGCAATTGAGACCA-TACTT-AGAAAACGGAAGACATATCGTTTCTGATATC
SEQ1417	AGCAGCTAATATTTCAAATATTTTAAACCCTGATTCTGTGGTTATT
SEQ1401_	TTGAATCTCTCA-AACATCGTTTGAGCCTCTATGGATTAACAAAAGATGACTTTCTCGG
SEQ1402	TTGAATCTCTCA-AACATCGTTTGAGCCTCTATGGATTAACAAAAGATGACTTTCTCGG
SEQ1403	TTGAATCTCTCA-AACATCGTTTGAGCCTCTATGGATTAACAAAAGATGACTTTCTCGG
SEQ1404	TTGAATCTCTCA-AACATCGTTTGAGCCTCTATGGATTAACAAAAGATGACTTTCTCGG
SEQ1405	GTGGCGGTGTCTCAGCAGCAGGTGAATTTTTACGTAGTCGCGTTGAGAAATACTTT
SEQ1406	TTGAATCTCTCA-AACATCGTTTGAGCCTCTATGGATTAACAAAAGATGACTTTCTCGG
SEQ1407	TTGAATCTCTCA-AACATCGTTTGAGCCTCTATGGATTAACAAAAGATGACTTTCTCGG
SEQ1408	GTGGCGGTGTCTCAGCAGCAGGTGAATTTTTACGTAGTCGCGTTGAGAAATACTTT
SEQ1409	TTGAATCTCTCA-AACATCGTTTGAGCCTCTATGGATTAACAAAAGATGACTTTCTCGG
SEQ1410	CAGCAGCAGCTGAATTTTTACGTAGTCGCGTTGAGAAATACTTT
SEQ1411	TTGAATCTCTCA-AACATCGTTTGAGCCTCTATGGATTAACAAAAGATGACTTTCTCGG
SEQ1412	GTGGCGGTGTCTCAGCAGCAGGTGAATTTTTACGTAGTCGCGTTGAGAAATACTTT
SEQ1413	TTGAATCTCTCA-AACATCGTTTGAGCCTCTATGGATTAACAAAAGATGACTTTCTCGG
SEQ1414	GTGGCGGTGTCTCAGCAGCAGGTGAATTTTTACGTAGTCGCGTTGAGAAATACTTT
SEQ1415	GTGGCGGTGTCTCAGCAGCAGGTGAATTTTTACGTAGTCGCGTTGAGAAATACTTT
SEQ1416	TTGAATCTCTCA-AACATCGTTTGAGCCTCTATGGATTAACAAAAGATGACTTTCTCGG
SEQ1417	GTGGCGGTGTCTCAGCAGCAGGTGAATTTTTACGTAGTCGCGTTGAGAAATACTTT
SEQ1401_	ATCGGTATGGGTTCTCCAGGAGCTGTTGATAGAACTAGTAAAACAGTAACAGGTGCTTT
SEQ1402	ATCGGTATGGGTTCTCCAGGAGCTGTTGATAGAACTAGTAAAACAGTAACAGGTGCTTT
SEQ1403	ATCGGTATGGGTTCTCCAGGAGCTGTTGATAGAACTAGTAAAACAGTAACAGGTGCTTT
SEQ1404	ATCGGTATGGGGTCTCCAGGAGCTGTTGATAGAACTAGTAAAAC
SEQ1405	GTCACATTTGCTTTCCCACAAGTTAAAAAGTCAACTA
SEQ1406	ATCGGTATGGGTTCTCCAGGAGCTG
SEQ1407	ATCGGTATGGGTTCTCCAGGAGCTGTTGATAGAACTAGTAAAACAGTAACAGGTGCTTT
SEQ1408	GTCACATTTGCTTTCCCACA
SEQ1409	ATCGGTATGGGTTCTCCAGGAGCTGTTGATAGAACTAGTAAAACAGTAACAGGTGCTTT
SEQ1410	GTCACATTTGCTTTCCCACAAGTTAAAAAGTCAACTAAAATTAAGATTGCTGAACTAGG
SEQ1411	ATCGGTATGGGTTCTCCAGGAGCTGTTGATAGAACTAGTAAAACAGTAACAGGTGCTTT
SEQ1412	GTCACATTTGCTTTCCCACAAGTTAAAAA
SEQ1413	ATCGGTATGGGTTCTCCAGGAGCTGTTGATAGAACTAGTAAAACAGTAACAGGTGCTTT
SEQ1414	ATCACATTTGCTTTCCCACAAGTTAAAAAGTCAACTAAAATTAAGATTG
SEQ1415	GTCACATTTGCTTTCCCACAAGTTAAAAAGTCAACTAA
SEQ1416	ATCGGTATGGGTTCTCCAGGAGCTGTTGATAGAACTAGTAAAACAGTCACAGGTGCTTT
SEQ1417	GTCACATTTGTTTTCCCACAAGGT



# Table 14: Comparative Sequences relating to SAG0471 (glucokinase)

SEQ1401_	AATCTAAATTGGGCTGATACTCAAGAAGTAGGTTCAGTTATTGAAAAAGAAGTTGGAAT
SEQ1402	AATCTAAATTGGGCTGATACTCAAGAAGTAGGTTCGGTTATTGAAAAAGAAGTTGGAAT
SEQ1403	AATCTAAATTGGGCTGATACTCAAGA
SEQ1404	A
SEQ1405	
SEQ1406	
SEQ1407	AATCTAAATTGGGCTGATACTCAAGAAGTAGGTTCAGTTATTGAAAAAGAAGTTGGAAT
SEQ1408	
SEQ1409	AATCTAAATTGGGCTGATACTCAAGAAGTAGGTTCGGTTATTGAAAAAGAAGTTGGAAT
SEQ1410	AATGAT
SEQ1411	AATCTAAATTGGGCTGATACTCAAGAAGTAGGTTCGGTTATTGAAAAAGAAGTTGGAAT
SEQ1412	
SEQ1413	AATCTAAATTGGGCTGATACTCAAGAAGTAGGTTCAGTTATTGAAAAAGAAGTTGGAAT
SEQ1414	
SEQ1415	
SEQ1416	AATCTAAATTGGGCTGATACTCAAGAAGTAGGTTCAGTTATTGAAAAAGAAGCTGGAAT
SEQ1417	**************************************
ocgiti,	
SEQ1401	CCA THURSDAN A HINCA THA A CCA THOCATA A HIGHWOOD A CCA THOCATA
SEQ1401_ SEQ1402	CCATTTTTTATTGATAACGATGCTAATGTTGCAGCACTTGGTGAACGCTGGGTAGGTGC
	CCATTTTTTTTTTTTTTTTTATTGATAACGATGCTAATGTTGCAGCACTTGGTGAACGCTGGGTAGGTGC
SEQ1403	**
SEQ1404	
SEQ1405	
SEQ1406 '	# 2 4 4 6 6 6 6 6 6 6 7 7 8 2 6 6 6 6 6 6 6 6 6 6 6 6 6 6 6 6 6 6
SEQ1407	CCATTTTTTATTGATAACGATGCTAATGTTGCAGCACTTGGTGAACGCTGGGTAGGTGC
SEQ1408	
SEQ1409	CCATTTTTTTTTTTATTGATAACGATGCTAATGTTGCAGCACTTGGTGAACGCTGGGTAGGTGC
SEQ1410	
SEQ1411	CCATTTTTTATTGATAACGATGCTAATGTTGCAGCACTTGGTGAACGCTGGGTAGGTGC
SEQ1412	
SEQ1413	CCATTTTTTATTGATAACGATGCTAATGTTGCAGCACTTGGTGAACGCTGGGTAGGTGC
SEQ1414	
SEQ1415	
SEQ1416	CCATTTTTTATTG
SEQ1417	
SEQ1401	
SEQ1402	GGTGCCAATAATCCCGACGTTGTTTTCGTAACCCTCGGAACAGGAGTAGGTGGAGGTGT GGTGCCAATAATCCCGATGTTGTTTTCGTAACCCTCGGAACAGGAGTAGGTGGAGGTGT
SEQ1403	GGTGCCAATAATCCCGATGTTGTTTTCGTAACCCTCGGAACAGGAGTAGGTGGAGGTGT
SEQ1403 SEQ1404	
SEQ1405	
SEQ1406	
SEQ1407	GGTGCCAATAATCCCGACGTTGTTTTCGTAACC
SEQ1408	
SEQ1409	GGTGCCAATAATCCCGATGTTGTTTTCGTAACCCTCGGAACAGGAGTAGGTGGAGGTGT
SEQ1410	
SEQ1411	GGTGCCAATAATCCCGATGTTGTTTTCGTAACCCTCGGAACAGGAGTA
SEQ1412	
SEQ1413	GGTGCCAATAATCCCGACGTTGTTTTCGTAACCCTCGGAACAGGAGTAGGTGGAGG
SEQ1414	
SEQ1415	
SEQ1416	
SEQ1417	

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# Table 14: C mparative Sequences relating to SAG0471 (glucokinase)

SEQ1401_	ATCGCAGATGGTAACCTCATCCATGGTGTTGCAGGAGCAGGTGGAGAAATTGGGCATAT
SEQ1402	ATCGCAGATGGTAACCTCATCCATGGTGTTGCAGGAGCAGGTGGAGAAATTGGGCATAT
SEQ1403	
SEQ1404	
SEQ1405	
SEQ1406	
SEQ1407	
SEQ1408	
SEQ1409	ATCGCAGATGGTAACCTCATCCATGGTGTTGCAAGAGCAGGTGGAGAAATTGGGCATAT
SEQ1410	
SEQ1411	
SEQ1412	
SEQ1413	
SEQ1414	
SEQ1415	
SEQ1416	
_	
SEQ1417	
0701404	
SEQ1401_	ATTGTTGATCCAGAAAATGGATTTACGTGCACATGTGGTAACAAAGGCTGCCTTGAGAC
SEQ1402	ATTGTTGATCCAGAKAATGGATTTACGTGCACATGTGGTAACAAAGGCTGTCTTGAGAC
SEQ1403	
SEQ1404	
SEQ1405	
SEQ1406	
SEQ1407	
SEQ1408	
SEQ1409	ATT
SEQ1410	
SEQ1411	
SEQ1412	
SEQ1413	
SEQ1414	
SEQ1415	
SEQ1416	
SEQ1417	
SEQ1401	GTTGCATCAGCGACAGGTGTTGTTAGAGTAGCACGTCAACTCGCAGAACAATATGAGGG
SEQ1402	GTTGCATCAGCGACAGGTGTTGTTAGAGTAGCACGTCAACTCGCAGAACAATATGAAGG
SEQ1403	
SEQ1404	
SEQ1405	
SEQ1406	
SEQ1407	
SEQ1407 SEQ1408	
SEQ1409	
SEQ1410	
SEQ1411	
SEQ1412	
SEQ1413	**
SEQ1414	
SEQ1415 .	6 - 6
SEQ1416	
SEQ1417	

# Table 14: Comparative Sequences relating t SAG0471 (glucokinase)

SEQ1401_	TCGTCTGCCATTAAAGCAGCGATTGACACCGGTGATACTGTTACAAGTAAAGATATTTT
SEQ1402	TCGTCTGCCATTAAAGCAGCGATTGACAACGGTGATACTGTTACAAGTAAAGATATTTT
SEQ1403	
SEQ1404	
SEQ1405	
SEQ1406	
SEQ1407	
SEQ1408	
SEQ1409	
SEQ1410	
SEQ1411	
SEQ1412	
SEQ1413	
SEQ1414	
SEQ1415	
SEQ1416	
SEQ1417	
ODQIAI!	
SEQ1401	ATAGCAGCAGAAGATGGGGATAAATTTGCTAATTCTGTTGTTGAACGTGTATCACGTTA
SEQ1402_	ATAGCAGCAGAAGATGGGGATAAATTTGCTAATTCTGTTGTTGAACGTGTATCACGTTA
SEQ1402	AIACCACACACACACACACACACACACACACACACACAC
SEQ1403	
-	
SEQ1405	
SEQ1406	
SEQ1407	
SEQ1408	
SEQ1409	
SEQ1410	
SEQ1411	
SEQ1412	
SEQ1413	
SEQ1414	
SEQ1415	
SEQ1416	
SEQ1417	
SEQ1401_	CTTGGACTGGCAGCAATATTTCAAATATTTTAAACCCTGATTCTGTGGTTATTGG
SEQ1402	CTTGGACTGGCAGCTAATATTTCAAATATTTTAAACCCTGATTCTGTGGTTATTGG
SEQ1403	4
SEQ1404	
SEQ1405	
SEQ1406	
SEQ1407	
SEQ1408	
SEQ1409	
SEQ1410	
SEQ1410 SEQ1411	
SEQ1412	
SEQ1412 SEQ1413	
SEQ1414	
SEQ1415	
SEQ1416	
SEQ1417	

# Table 14: C mparative Sequences relating to SAG0471 (glucokinase)

SEQ1401	
OCCIACI	GGCGGTGTCTCAGCAGCAGGTGAATTTTTACGTAGTCGCGTTGAGAAATACTTTGTCAC
SEQ1402	GGCGGTGTCTCAGCAGCAGGTGAATTTTTACGTAGTCGCGTTGAGAAATACTTTGTCAC
SEQ1403	_4
SEQ1404	B
SEQ1405	
SEQ1406	
SEQ1407	
SEQ1408	
SEQ1409	
SEQ1410	
SEQ1411	0
SEQ1412	, 
SEQ1413	
SEQ1414	#FEVU9988###8##=
SEQ1415	
SEQ1416	
SEQ1417	
SEQ1401_	TTTGCTTTCCCACAAGTTAAAAAGTCAACTAAAATTAAGAT
SEQ1402	TTTG
SEQ1403	
SEQ1404	***********
SEQ1404 SEQ1405	
SEQ1405	
SEQ1405 SEQ1406	
SEQ1405 SEQ1406 SEQ1407	
SEQ1405 SEQ1406 SEQ1407 SEQ1408	
SEQ1405 SEQ1406 SEQ1407 SEQ1408 SEQ1409	
SEQ1405 SEQ1406 SEQ1407 SEQ1408 SEQ1409 SEQ1410	
SEQ1405 SEQ1406 SEQ1407 SEQ1408 SEQ1409 SEQ1410 SEQ1411	
SEQ1405 SEQ1406 SEQ1407 SEQ1408 SEQ1409 SEQ1410 SEQ1411 SEQ1412	
SEQ1405 SEQ1406 SEQ1407 SEQ1408 SEQ1409 SEQ1410 SEQ1411 SEQ1412 SEQ1413	
SEQ1405 SEQ1406 SEQ1407 SEQ1408 SEQ1410 SEQ1411 SEQ1411 SEQ1411 SEQ1412 SEQ1413 SEQ1414	



#### Table 15: Comparative Sequences relating to SAG0492

SEQ ID NO. 1506: SAG0492 FROM THE A909 GBS TYPE IA STRAIN
CAATACAAGGACTTCATAAAAGTTTTGGGAAAAATGAGGTTTTAAAAGGCATTGACTTGGATATTCATCAAGGAGAAGTAGTGGTTATT
ATTGGCCCTTCTGGCTCTGGTAAGTCAACATTTTTAAGAACAATGAATCTCTTGGAAGTACCAACAAAGGGAACAGTGACTTTTGAAGG
GATTGATATAACAGACAAAAAGAATGATATTTTTAAAATGCGCGAAAAAATGGGCATGGTTTTTCAACAGTTCAATCTATTTCCCAATA
TGACTGTACTTGAATATTACTTTATCACCTATTAAGACAAAGGGGCTTTCTTAAGTTGCACAACAACAACAAAAGCATATGAAGTCT
GAAAAAGTTGGACTCAAAGAGAAGGCTAATACTTATCCAGCTCAGCTTATCTGAGGACAACAACAACAACAATTGCTATTGCAAGAATTGCAAGAATTGCTATTTTTATGCAAGAATTCCTGAATCGTAATCTGGATGTCATTTTTATGCAAG
ATTTAGCTAAATCTGGTATGACGATGGTTATTTGCACTCATGAAATGGTTTTTGCACTGTAAGCGATTGCTTTATGGAT
GCAGGAATTATTGTGAGCAAGGGGCCCCTAAGGAAGTATTTTGAGCAGACAAAAGAAATCCGCACAAAGAGATTTCTT

#### Table 15: Comparative Sequences relating to SAG0492

SEQ ID NO. 1508: SAG0492 FROM THE H36b GBS TYPE ID STRAIN
ATGAGGTTTTAAAAAGGCATTGACTTGGATATTCATCAAGGAGAAGTAGTGGTTATTATTGGCCCTTCTGGCTCTGGTAAGTCAACATTT
TTAAGAACAATGAATCTCTTGGAAGTACCAACAAAAGGGAACAGTGACTTTTGAAGGGATTGATATAACAGACAAAAAGAATGATATTTT
TAAAATGCGCGAAAAAATGGGCATGGTTTTTCAACAGTTCAATCTATTTCCCAATATGACTGTACTAGAAAATATTACTTTATCACCTA
TTAAGACAAAGGGGCTTTCTAAGCTTGATGCTCAGACAAAAGCATATGAGCTACTTGAAAAAAGTTGGACTCAAAGAGAAGGCTAATACT
TATCCAGCTAGCTTATCTGGAGGACAACAACAACGAATTGCTATTGCAAGAGTCTTGCAATGAATCCTGATGACCTTCTTTTTTGATGA
ACCTACTTCAGCTCTTGATCCTGAAATGGTAAGGTGAAGTCTTGACTGTATTGCAAGATTTAGCTAAATCTGGTATGACGATGGTTATTG
TCACTCATGAAATGGGTTTTGCACGTGAAGTAGCGGATCGTTCATTTTTATGGATGCASGAATTATTGTTGAGCAAGGGGCCCCCTAAG
GAAGTAT

SEQ ID NO. 1510: SAG0492 FROM THE M732 GBS TYPE III STRAIN
GGTGGTTATTATTGGCCCTTCTGGCTCTGGTAAGTCAACATTTTTAAGAACAATGAATCTCTTGGAAGTACCAACAAAGGGAACAGTGA
CTTTTGAAGGGATTGATATAACAGACAAAAAGAATGATATTTTTAAAATGCGCGAAAAAATGGGCATGGTTTTTCAACAGTTCAATCTA
TTTCCCAATATGACTGTACTAGAAAAATATTACTTTATCACCTATTAAGACAAAGGGACTTTCTAAGCTTGATGCTCAGACAAAAGCATA
CGAGCTACTTGAAAAAAGTTGGACTCAAAGAGAAAGGCTAATGCTTATCCAGCAAGCTTATCTGG

SEQ ID NO. 1511: SAG0492 FROM THE COHI GBS TYPE IA STRAIN
ATTGACTTGGATATTCATCAAGGAGAAGTGGTTGTTATTATTGGCCCTTCTGGCTCTGGTAAGTCAACATTTTTAAGAACAATGAATCT
CTTGGAAGTACCAACAAAGGGAACAGTGACTTTTGAAGGGATTGATATAACAGACAAAAAGAATGATTTTTAAAATGCGCGAAAAAA
TGGGCATGGTTTTTCAACAGTTCAATCTATTTCCCAATATGACTGTACTAGAAAATATTACTTTATCACCTATTAAGACAAAGGGACTT
TCTAAGCTTGATGCTCAGACAAAAGCATACGAGCTACTTGAAAAAGTTGGACTCAAAGAGAAGGCTAATGCTTATCCAGCAAGCTTATC
TGG

## Table 15: Comparative Sequences relating t SAG0492

SEQ1501	TGACTTGG
SEQ1502	TGACTTGG TGACTTGACT
SEQ1503	AAAATGAGGTTTTAAAAGGCATTGACTTGG
SEQ1504	GAGGTTTTAAAAGGCATTGACTTGC
SEQ1505	
SEQ1506	AATACAAGGACTTCATAAAAGTTTTGGGAAAAATGAGGTTTTAAAAGGCATTGACTTGG
SEQ1507	
SEQ1508	GACTTGGATGAGGTTTTAAAAGGCATTGACTTGGGGTTTTAAAAGGCATTGACTTGG
SEQ1509	ATGAGGTTTTAAAAGGCATTGACTTGG
SEQ1510	GGTTTTAAAAGGCATTGACTTGG
SEQ1511	ATTGACTTGG
2EQ1211	ATTGACTTGG
GT01 F01	
SEQ1501	TATTCATCAAGGAGAGTGGTGGTTATTATTGGCCCTTCTGGCTCTGGTAAGTCAACAT
SEQ1502	TATTCATCAAGGAGAAGTAGTGGTTATTATTGGCCCTTCTGGCTCTGGTAAGTCAACAT
SEQ1503	TATTCATCAAGGAGAAGTAGTGGTTATTATTGGCCCTTCTGGCTCTGGTAAGTCAACAT
SEQ1504	TATTCATCAAGGAGAAGTGGTGGTTATTATTGGCCCCTTCTGGCTCTGGTAAGTCAACAT
SEQ1505	TGGTGGTTATTATTGGCCCTTCTGGCTCTGGTAAGTCAACAT
SEQ1506	TATTCATCAAGGAGAAGTAGTGGTTATTATTGGCCCCTTCTGGCTCTGGTAAGTCAACAT
SEQ1507	TATTCATCAAGGAGAAGTGGTGGTTATTATTGGCCCCTTCTGGCTCTGGTAAGTCAACAT
SEQ1508	TATTCATCAAGGAGAAGTAGTGGTTATTATTGGCCCTTCTGGCTCTGGTAAGTCAACAT
SEQ1509	TATTCATCAAGGAGAAGTAGTGGTTATTATTGGCCCCTTCTGGCTCTGGTAAGTCAACAT
SEQ1510	GGTGGTTATTATTGGCCCTTCTGGCTCTGGTAAGTCAACAT
SEQ1511	TATTCATCAAGGAGAAGTGGTGGTTATTATTGGCCCTTCTGGCTCTGGTAAGTCAACAT
SEQ1501	TTTTAAGAACAATGAATCTCTTGGAAGTACCAACAAGGGAACAGTGACTTTTGAAGGAA
SEQ1502	TTTTAAGAACAATGAATCTCTTGGAAGTACCAACAAAGGGAACAGTGACTTTTGAAGGGA
SEQ1503	TTTTAAGAACAATGAATCTCTTGGAAGTACCAACAAAGGGAACAGTGACTTTTGAAGGGA TTTTAAGAACAATGAATCTCTTGGAAGTACCAACAAAGGGAACAGTGACTTTTGAAGGGA
SEQ1504	TTTTAAGAACAATGAATCTCTTGGAAGTACCAACAAAGGGAACAGTGACTTTTGAAGGGA TTTTAAGAACAATGAATCTCTTGGAAGTACCAACAAAGGGAACAGTGACTTTTGAAGGGA
SEQ1505	TITIAAGAACAATGAATCICTTGGAAGTACCAACAAAGGGAACAGTGACTTTTGAAGGGA
SEQ1506	TTTTAAGAACAATGAATCTCTTGGAAGTACCAACAAAGGGAACAGTGACTTTTGAAGGGA
	TTTTAAGAACAATGAATCTCTTGGAAGTACCAACAAAGGGAACAGTGACTTTTGAAGGGA
SEQ1507	TTTTAAGAACAATGAATCTCTTGGAAGTACCAACAAAGGGAACAGTGACTTTTGAAGGGA
SEQ1508	TTTTAAGAACAATGAATCTCTTGGAAGTACCAACAAAGGGAACAGTGACTTTTGAAGGGA
SEQ1509	TTTTAAGAACAATGAATCTCTTGGAAGTACCAACAAAGGGAACAGTGACTTTTGAAGGGA
SEQ1510	TTTTAAGAACAATGAATCTCTTGGAAGTACCAACAAAGGGAACAGTGACTTTTGAAGGGA
SEQ1511	TTTTAAGAACAATGAATCTCTTGGAAGTACCAACAAAGGGAACAGTGACTTTTGAAGGGA
SEQ1501	TTGATATAACAGACAAAAAAATGATATTTTTAAAATGCGCGAAAAAATGGGCATGGTTT
SEQ1502	TTGATATAACAGACAAAAAGAATGATATTTTTAAAATGCGCGAAAAAATGGGCATGGTTT
SEQ1503	TTGATATAACAGACAAAAAGAATGATATTTTTAAAATGCGCGAAAAAATGGGCATGGTTT
SEQ1504	TTGATATAACAGACAAAAAGAATGATATTTTTAAAATGCGCGAAAAAATGGGCATGGTTT
SEQ1505	TTGATATAACAGACAAAAAGAATGATATTTTTAAAATGCGCGAAAAAATGGGCATGGTTT
SEQ1506	TTGATATAACAGACAAAAAGAATGATATTTTTAAAATGCGCGAAAAAATGGGCATGGTTT
SEQ1507	TTGATATAACAGACAAAAAGAATGATATTTTTAAAATGCGCGAAAAAATGGGCATGGTTT
SEQ1508	TTGATATAACAGACAAAAAGAATGATATTTTTAAAATGCGCGAAAAAATGGGCATGGTTT
SEQ1509	TTGATATAACAGACAAAAAGAATGATATTTTTAAAATGCGCGAAAAAATGGGCATGGTTT
SEQ1510	TTGATATAACAGACAAAAAGAATGATATTTTTAAAATGCGCGAAAAAATGGGCATGGTTT
SEQ1511	TTGATATAACAGACAAAAAGAATGATATTTTTAAAATGCGCGAAAAAATGGGCATGGTTT
022221	11GATATAACAGACAAAAAGATATTTTTAAAATGCGCGAAAAAATGGGCATGGTTT
SEQ1501	
SEQ1501 SEQ1502	TTCAACAGTTCAATCTATTTCCCAATATGACTGTACTAGAAAATATTACTTTATCACCTA
SEQ1502 SEQ1503	TTCAACAGTTCAATCTATTTCCCAATATGACTGTACTAGAAAATATTACTTTATCACCTA
	TTCAACAGTTCAATCTATTTCCCAATATGACTGTACTAGAAAATATTACTTTATCACCTA
SEQ1504	TTCAACAGTTCAATCTATTTCCCAATATGACTGTACTAGAAAATATTACTTTATCACCTA
SEQ1505	TTCAACAGTTCAATCTATTTCCCAATATGACTGTACTAGAAAATATTACTTTATCACCTA
SEQ1506	TTCAACAGTTCAATCTATTTCCCAATATGACTGTACTAGAAAATATTACTTTATCACCTA
SEQ1507	TTCAACAGTTCAATCTATTTCCCAATATGACTGTACTAGAAAATATTACTTTATCACCTA
SEQ1508	TTCAACAGTTCAATCTATTTCCCAATATGACTGTACTAGAAAATATTACTTTATCACCTA
SEQ1509	TTCAACAGTTCAATCTATTTCCCAATATGACTGTACTAGAAAATATTACTTTATCACCTA
SEQ1510	TTCAACAGTTCAATCTATTTCCCAATATGACTGTACTAGAAAATATTACTTTATCACCTA
SEQ1511	TTCAACAGTTCAATCTATTTCCCAATATGACTGTACTAGAAAATATTACTTTATCACCTA
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## Table 15: Comparative Sequences relating to SAG0492

SEQ1501	TTAAGACAAAGGGACTTTCTAAGCTTGATGCTCAGACAAAAGCATACGAGCTACTTGAAA
SEQ1502	TTAAGACAAAGGGGCTTTCTAATCTTGATGCTCAGACAAAAGCATATGAGCTACTTGAAA
SEQ1503	TTAAGACAAAGGGGCTTTCTAATCTTGATGCTCAGACAAAAGCATATGAGCTACTTGAAA
SEQ1504	TTAAGACAAAGGGACTTCTAAGCTTGATGCTCAGACAAAAGCATACGAGCTACTTGAAA
<del></del>	TIALGACAAAGGACTTTCTAAGCTTGATGCTCAGACAAAAGCATACGAGCTACTTGAAA
SEQ1505	TTAAGACAAAGGGACTTTCTAAGCTTGATGCTCAGACAAAAGCATACGAGCTACTTGAAA
SEQ1506	TTAAGACAAAGGGGCTTTCTAAGCTTGATGCTCAGACAAAAGCATATGAGCTACTTGAAA
SEQ1507	TTAAGACAAAGGGACTTTCTAAGCTTGATGCTCAGACAAAAGCATACGAGCTACTTGAAA
SEQ1508	TTAAGACAAAGGGGCTTTCTAAGCTTGATGCTCAGACAAAAGCATATGAGCTACTTGAAA
SEQ1509	TTAAGACAAAGGGGCTTTCTAAGCTTGATGCTCAGACAAAAGCATATGAGCTACTTGAAA
SEQ1510	TTAAGACAAAGGGACTTTCTAAGCTTGATGCTCAGACAAAAGCATACGAGCTACTTGAAA
SEO1511	I A CO CA A COCA COMMON A COMMON A COLOR COLOR CA
SPOISIT	TTAAGACAAAGGGACTTTCTAAGCTTGATGCTCAGACAAAAGCATACGAGCTACTTGAAA
CEO1 501	A A CHIMCOLOGICAL AND A A CALLOGODIA A COMPANION AND A CALLOGODIA A CA
SEQ1501	AAGTTGGACTCAAAGAGAGAGGCTAATGCTTATCCAGCTAGCT
SEQ1502	AAGTTGGACTCAAAGAGAGAGGCTAATACTTATCCAGCTAGCT
SEQ1503	AAGTTGGACTCAAAGAGAAGGCTAATACTTATCCAGCTAGCT
SEQ1504	AAGTTGGACTCAAAGAGGAGGCTAATGCTTATCCAGCAAGCTTATCTGGAGGACAACAAC
SEQ1505	AAGTTGGACTCAAAGAGAAGGCTAATGCTTATCCAGCTAGCT
SEQ1506	AAGTTGGACTCAAAGAGAAGGCTAATACTTATCCAGCTAGCT
SEQ1507	AAGTTGGACTCAAAGAGGCTAATGCTTATCCAGCTAGCTTATCTGGAGGACAACAAC
SEQ1508	AAGTTGGACTCAAAGAGGCTAATACTTATCCAGCTAGCTTATCTGGAGGACAACAAC
SEQ1509	
	AAGTTGGACTCAAAGAGAAGGCTAATACTTATCCAGCTAGCT
SEQ1510	AAGTTGGACTCAAAGAGAAGGCTAATGCTTATCCAGCAAGCTTATCTGG
SEQ1511	AAGTTGGACTCAAAGAGAAGGCTAATGCTTATCCAGCAAGCTTATCTGGTABCMARATVS
SEQ1501 -	ACGGATTGCTATTGCAAGAGGTCTTGCAATGAATCCTGATGTCCTTCTTTTTGATGAAC
SEQ1502	ACGAATTGCTATTGCAAGAGGTCTTGCAATGAATCCTCATGTCCTTCTTTTTGATGAAC
SEQ1503	ACGAATTGCTATTGCAAGAGGTCTTGCAATGAATCCTGATGTCCTTCTTTTTGATGAAC
SEQ1504	ACGGATTGCTATTGCAAGAGGTCTTGCAATGAATCCTGATGTCCTTCTTTTTGATGAAC
SEQ1505	ACGAATTGCTATTGCAAGAGGTCTTGCAATGAATCCTGATGTCCTTCTTTTTGATGAAC
SEQ1506	ACGAATTGCTATTGCAAGAGGTCTTGCAATGAATCCTGATGTCCTTCTTTTTGATGAAC
SEQ1507	ACGAATTGCTATTGCAAGAGGTCTTGCAATGAATCCTGATGTCCTTCTTTTTGATGAAC
SEQ1508	
SEQ1509	ACGAATTGCTATTGCAAGAGGTCTTGCAATGAATCCTGATGTCCTTCTTTTTGATGAAC
	ACGAATTGCTATTGCAAGAGGTCTTGCAATGAATCCTGATGTCCTTCTTTTTGATGAAC
SEQ1510	
SEQ1511	NCSRATNGTSAG
SEQ1501	TACTTCAGCTCTTGATCCTGAAATGGTAGGTGAAGTCTTGACTGTTATGCAAGATTTAG
SEQ1502	TACTTCAGCTCTTGATCCTGAAATGGTAGGTGAAGTCTTGACTGTTATGCAAGATTTAG
SEQ1503	TACTTCAGCTCTTGATCCTGAAATGGTAGGTGAAGTCTTGACTGTTATGCAAGATTTAG
SEQ1504	TACTTCAGCTCTTGATCCTGAAATGGTAGGTGAAGTCTTGACTGTTATGCAAGATTTAG
SEQ1505	TACTTCAGCTCTTGATCCTGAAATGGTAGGTGAAGTCTTGACTGTTATGCAAGATTTAG
SEQ1506	TACTTCAGCTCTTGATCCTGAAATGGTAGGTGAAGTCTTGACTGTTATGCAAGATTTAG
SEQ1507	TACTTCAGCTCTTGATCCTGAAATGGTAGGTGAAGTCTTGACTGTTATGCAAGATTTAG
SEQ1508	TACTTCAGCTCTTGATCCTGAAATGGTAGGTGAAGTCTTGACTGTTATGCAAGATTTAG
_	
SEQ1509	TACTTCAGCTCTTGATCCTGAAATGGTAGGTGAAGTCTTGACTGTTATGCAAGATTTAG
SEQ1510	
SEQ1511	
SEQ1501	TAAATCTGGTATGACGATGGTTATTGTCACTCATGAAATGGGTTTTGCACGTGAAGTAG
SEQ1502	TAAATCTGGTATGACGATGGTTATTGTCACTCATGAAATGGGTTTTGCACGTGAAGTAG
SEQ1503	TAAATCTGGTATGACGATGGTTATTGTCACTCATGAAATGGGTTTTGCACGTGAAGTAG
SEQ1504	TAAATCTGGTATGACGATGGTTATTGTCACTCATGAAATGGGTTTTGCACGTGAAGTAG
SEQ1505	TAAATCTGGTATGACGATGGTTATTGTCACTCATGAAATGGGTTTTGCACGTGAAGTAG
SEQ1506	TAAATCTGGTATGACGATGGTTATTGTCACTCATGAAATGGGTTTTGCACGTGAAGTAG
SEQ1507	TAAATCTGGTATGACGATGGTTATTGTCACTCATGAAATGGGTTTTGCACGTGAAGTAG
SEQ1507	
· · · · · · · · · · · · · · · · · · ·	TAAATCTGGTATGACGATGGTTATTGTCACTCATGAAATGGGTTTTGCACGTGAAGTAG
SEQ1509	TAAATCTGGTATGACGATGGTTATTGTCACTCATGAAATGGGTTTTGCACGTGAAGTAG
SEQ1510	
SEQ1511	~

#### 50406 7.082602

# Table 15: Comparative Sequences relating t SAG0492

SEQ1501	GGATCGTGTCATTTTATGGATGCAGGCATTATTGT-GAGCAAGGGACCCCTAAGGAAG
SEQ1502	GGATCGTGTCATTTTTATGGACGCAGAAATTAT
SEQ1503	GGATCGTGTCATTTTATGGATGCAGGAATTATTGTTGAGCAAGGGGCCC
SEQ1504	GGATCGTGTCATTTTATGGATGCAGGGATTATTGTTGAGCAAGGGACCCCTAAGAAAG
SEQ1505	GGATCGTGTCATTTTATGGATGCAGGCATTATTGTTGASCAAGGGACCCCTAAGGAAG
SEQ1506	GGATCGTGTCATTTTTATGGATGCAGGAATTATTGTGAGCAAGGGGCCCCTAAGGAAGT
SEQ1507	GGATCGTGTC-TTTTTATGGATGCGGGAATTATTGT-GAGCAAGGGACC
SEQ1508	GGATCGTGTCATTTTATGGATGCASGAATTATTGTTGAGCAAGGGGCCCCTAAGGAAG
SEQ1509	GGATCGTGTCATTTTATGGATGCAGGAATTATTGTTGAGCAAGGGGCCCCTAAGGAAG
SEQ1510	GOATCOTOTOTITITATOGATGCAGGAATTATTGTTGAGCAAGGGGCCCCTAAGGAAG
SEQ1511	
SEQ1501	AT
SEQ1502	
SEQ1503	
SEQ1504	AT
SEQ1505	A
SEQ1506	TTTGAGCAGACAAAAGAAATCCGCACAAGAGATTTCTT
SEQ1507	111 discrimination Code Control 11011
SEQ1508	AT
SEQ1509	ATTTAGCAAAACAAAGAAAT
SEQ1510	AT I TOURAGAMAGAMAI
SEQ1510 SEO1511	
GEVICIT	

#### Sable 16: Comparative Sequences relating to SAG0767 (D-alanine – D-alanine ligase)

SEQ ID NO. 1604: SAG0767 FROM THE CJB110 GBS NONTYPEABLE STRAIN REVERSE COMPLEMENT

CGTCGATTTCTATGACTATGACGCCAAATATATTGATAATAAAATTACTATGGATATTCCAGCTAAAGTTGATGAAGCAA CTATGGAAGCAATGCGTCAATATGCAAGTAAAGCTTTTAAAGCAATCGGGGCTTGTGGTTTATCACGCTGTGATTTCTTT TTGACGAAAGATGGACAAATCTTCTTAAACGAACTGAATACAATGCCC

- SEQ ID NO. 1606: SAG0767 FROM THE 1169NT1 GBS TYPE V STRAIN REVERSE COMPLEMENT CTAATATGGGGTCATCAGTAGGTATTTCAAAAGCGACAGATGAATCCTCACTTCGCTCTGCAATTGACTTAGCTCTCAAG TATGATAGCCGTATTTTGATTGAACAAGGCGTGACAGCTCGTGAAATTGAAGTAGGTATTTTAGGCAATAATGATGTTAA GACAACTTTTCCTGGCGAAGTCGTTAAAGACGTCGATTTCTATGACTATGACGCCAAATATATTGATAATAAAATTACTA TGGATATTCCAGCTAAAGTTGATGAAGCAACTATGGAAGCAATGCGTCAATATGCAAGTAAAGCTTTTAAAGCAATCGGG GCTTGTGGTTTATCACGCTGTGATTTCTTTTTGACGAAAGATGGACAAATCTTCTTAAACGAACTGAATACAATGCCCGG TTTTACTCAGTGGTCAATGTATCCTCTGCTTTTGGGGAAAAT



#### Table 16: Comparative Sequences relating to SAG0767 (D-alanine – D-alanine ligase)

SEQ ID NO. 1613: SAG0767 FROM THE M732 GBS TYPE III STRAIN REVERSE COMPLEMENT ATGCGATTAAACTCTCTTTAGAACCTTTAAGTTTCCCAATTTTTGTAAACCCGGCTAATATGGGGTCATCAGTAGGTATT TCAAAAGCGACAGATGAATCCTCACTTCGCTCTGCAATTGACTTAGCTCTCAAGTATGATAGCCGTATTTTGATCAACA AGGCGTGACAGCTCGTGAAATTGAAGTAGGTATTTTAAGCCAATAATGATGTTAAGACAACTTTTCCTGGCGAAGTTGTTA AAGACGTCGATTTCTATGACTATGACGCCAAATATATTGATAATAAAATTACTATGGATATTCCAGCTAAAGTTGATGAA GCAACTATGGAAGCAATGCGTCAATATGCAAGTAAAGCTTTTAAAGCAATCGGGGCTTGTGGTTTATCACGCTGTGATTT CTTTTTGACGAAAGATGGACAAATCTTCTTAAACGAACTGAATACAATGCCCGGTTTTACTCAGTGGTCAATGTATCCTC TGCTTTGGGAAAATATTGGGGCTAACTT

SEQ ID NO. 1615: SAG0767 FROM THE A909 GBS TYPE IA STRAIN REVERSE COMPLEMENT TTTTGAGGGTGATGATTTGGAACATGCGATTAAACTCTCTTTAGAAACTTTTAAGTTTCCCAATTTTTGTAAAACCGGCTA ATATGGGGTCATCAGTAGTTTCCAAAAGCGACAGATGAATCCTCACTTCGCTCTGCAATTGACTTAGCTCTCAAGTAT GATAGCCGTATTTTGATTGAACAAGGCGTGACAGCTCGTGAAATTGAAGTAGGTATTTTAGGCCAATAATGATGTTAAGAC AACTTTTCCTGGCGAAGTCGTTAAAGACGTCGATTTCTATGACGAACTATGACGCCCAAATATATTGATAAAAATTACTATGG ATATTCCAGCTAAAGTTGATGAAGCAACTATGGAAGCAATGCGTCAATATGCAAGTAAAGCTTTTAAAACAATCCGGGGCT TGTGGTTTATCACGCTGTGATTTCTTTTTGACGAAAGATGGACAAATCTTCTTAAACGAACTGAATACAATGCCCGGTTT TACTCAGTGGTCAATGTTATCCCCTGCTTTTGGGAAAATATTGGGGCTAACTTATAGTGA

## Table 16: Comparative Sequences relating to SAG0767 (D-alanine - D-alanine ligase)

SEQ ID NO. 1617: SAG0767 FROM THE JM9130013 GBS TYPE VIII STRAIN REVERSE COMPLEMENT

AAGCAGGGGATACATTGACCACTGAGTAAAACCGGGCATTGTATTCAGTTCGTTTAAGAAGATCTGTCCATCTTTCGTCA
AAAAGAAATCACAGCGTGATAAACCACAAGCCCCGATTGCTTTAAAAGCTTTACTTGCATATTGACGCATTGCTTCCATA
GATGCTTCATCAACTTTAGCTGGAATATCCATAGCAATTTTATTATCAATATTTTGGCG

SEQ1601	GGTCGCTCTGTCGGAACGTGAAGTATCTGTACTGTCTGCAGAAAGCGTCATGCGTGCTA
SEQ1602	
SEQ1603	#=+= ***********************************
SEQ1604	
SEQ1605	
SEQ1606	
SEQ1607	
SEQ1608	
SEQ1609	
SEQ1610	75648455554744558845884478884578845788888888
SEQ1611	
SEQ1612	
SEQ1613	
SEQ1614	
SEQ1615	
SEQ1616	
SEQ1617	
SEQ1601	TAATTATGATAAATTTTTTGTTAAAACTTATTTTATCACGCAAGTAGGTCAATTTATTA
SEQ1602	THE CATACAT THE TATACAT TATTA
SEQ1603	
SEQ1604	
SEQ1605	
SEQ1605	
SEQ1607	
SEQ1608	
SEQ1609	
SEQ1610	
SEQ1611	
SEQ1612	
SEQ1613	
SEQ1614	
SEQ1615	~
SEQ1616	
SEQ1617	
SEQ1601	AACACAAGAATTTGATGAAATGCCATCTTCAGATGAAAAGTTAATGACAAACCAAACTG
SEQ1602	
SEQ1603	
SEQ1604	
SEQ1605	
SEQ1606	
SEQ1607	
SEQ1608	
SEQ1609	
SEQ1610	~==~==================================
SEQ1611	
SEQ1612	
SEQ1613	
SEQ1614	
SEQ1615	*
SEQ1616	
SEQ1617	~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~

## Table 16: Comparative Sequences relating to SAG0767 (D-alanine – D-alanine ligase)

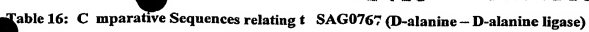
SEQ1601	TGATTTAGACAAAATGGTTCGTCCAAGTGATATCTATGATGATAATGCAATTGTTTTCC
SEQ1602	
SEQ1603	
SEQ1604	
SEQ1605	
SEQ1606	
SEQ1607	,
SEQ1608	
SEQ1609	
SEQ1610	400044000000000000000000000000000000000
SEQ1611	# 0 = 0 = 4 = 4 = 0 = 4 = 0 = 0 = 4 = 4 =
SEQ1612	
SEQ1613	
SEQ1614	*
SEQ1615	
SEQ1616	**
SEQ1617	
_	
SEQ1601	CGTTTTACATGGACCAATGGGGAAGATGGTTCTATCCAAGGATTTTTAGAAGTTTTAA
SEQ1602	
SEQ1603	
SEQ1604	
SEQ1605	**************************************
SEQ1606	
SEQ1607	000000
SEQ1608	
SEQ1609	
SEQ1610	
SEQ1611	
SEQ1612	
SEQ1613	
SEQ1614	
SEQ1615	
SEQ1616	*
SEQ1617	
SEQ1601	GATGCCTTATGTTGGGACTAATATTCTATCTTCAAGCGTGGCTATGGATAAAATTACAA
SEQ1602	
SEQ1603	4
SEQ1604	
. SEQ1605	
SEQ1606	
SEQ1607	
SEO1608	**************************************
SEQ1609	GGCTATGGATAAATTACAA
SEQ1610	
SEQ1611	
SEQ1612	*
SEQ1613	
SEQ1614	
SEO1615	
SEQ1616 SEQ1616	
SEQ1617	
PRÄTOT,	

# Table 16: Comparative Sequences relating to SAG0767 (D-alanine – D-alanine ligase)

SEQ1601	AAAACAAGTCCTTGCAACAGTAGGTGTACCTCAGGTTGCATATCAAACTTATTTTGAGG
SEQ1602	
SEQ1603	
SEQ1604	
SEQ1605	
SEQ1606	***************************************
SEQ1607	
SEQ1608	
SEQ1609	AAAACAAGTCCTTGCAACAGTAGGTGTACCTCAGGTTGCATATCAAACTTATTTTGAGG
SEQ1610	
SEQ1611	
SEQ1612	
SEQ1613	~
SEQ1614	
SEQ1615	TTTTGAGG
SEQ1616	
SEQ1617	
• • • • • • • • • • • • • • • • • • • •	
SEQ1601	TGATGATTTGGAACATGCGATTAAACTCTCTTTAGAAACTTTAAGTTTCCCAATTTTTG
SEQ1602	
SEQ1603	
SEQ1604	~
SEQ1605	*
SEQ1606	*
SEQ1607	*
SEQ1608	
SEQ1609	TGATGATTTGGAACATGCGATTAAACTCTCTTTAGAAACTTTAAGTTTCCCAATTTTTG
SEO1610	
SEQ1611	
SEQ1612	
SEQ1613	ATGCGATTAAACTCTCTTTAGAACCTTTAAGTTTCCCAATTTTTG
SEQ1614	THE CONTINUE OF THE CONTINUE O
SEQ1615	
SEQ1615	TGATGATTTGGAACATGCGATTAAACTCTCTTTAGAAACTTTAAGTTTCCCAATTTTTG
_	
SEQ1615 SEQ1616	
SEQ1615 SEQ1616	TGATGATTTGGAACATGCGATTAAACTCTCTTTAGAAACTTTAAGTTTCCCAATTTTTG
SEQ1615 SEQ1616 SEQ1617	TGATGATTTGGAACATGCGATTAAACTCTCTTTAGAAACTTTAAGTTTCCCAATTTTTG
SEQ1615 SEQ1616 SEQ1617 SEQ1601	TGATGATTTGGAACATGCGATTAAACTCTCTTTAGAAACTTTAAGTTTCCCAATTTTTG
SEQ1615 SEQ1616 SEQ1601 SEQ1602	TGATGATTTGGAACATGCGATTAAACTCTCTTTAGAAACTTTAAGTTTCCCAATTTTTG
SEQ1615 SEQ1616 SEQ1617 SEQ1601 SEQ1602 SEQ1603	AAAACCGGCTAATATGGGGTCATCAGTAGGTATTTCAAAAGCGACAGATGAATCCTCAC
SEQ1615 SEQ1616 SEQ1617 SEQ1601 SEQ1602 SEQ1603 SEQ1604	TGATGATTTGGAACATGCGATTAAACTCTCTTTAGAAACTTTAAGTTTCCCAATTTTTG
SEQ1615 SEQ1616 SEQ1617 SEQ1601 SEQ1602 SEQ1603 SEQ1604 SEQ1605	TGATGATTTGGAACATGCGATTAAACTCTCTTTAGAAACTTTAAGTTTCCCAATTTTTG  AAAACCGGCTAATATGGGGTCATCAGTAGGTATTTCAAAAGCGACAGATGAATCCTCACAAACCGGGCTCGCTCTGCGGAACGTGAAGTATCTGTACTG-TCTGCAGAAA-GCGTAACGTGAAGTATCTGTACTGCTCTGCAGAAAAGCGT
SEQ1615 SEQ1616 SEQ1617 SEQ1601 SEQ1602 SEQ1603 SEQ1604 SEQ1605 SEQ1606	TGATGATTTGGAACATGCGATTAAACTCTCTTTAGAAACTTTAAGTTTCCCAATTTTTG  AAAACCGGCTAATATGGGGTCATCAGTAGGTATTTCAAAAGCGACAGATGAATCCTCACAACGTGAAGTATCTGTACTG-TCTGCAGAAA-GCGT
SEQ1615 SEQ1616 SEQ1617 SEQ1601 SEQ1602 SEQ1603 SEQ1604 SEQ1605 SEQ1606 SEQ1606	TGATGATTTGGAACATGCGATTAAACTCTCTTTAGAAACTTTAAGTTTCCCAATTTTTG  AAAACCGGCTAATATGGGGTCATCAGTAGGTATTTCAAAAGCGACAGATGAATCCTCACAAACCGGGC
SEQ1615 SEQ1616 SEQ1617 SEQ1601 SEQ1602 SEQ1603 SEQ1604 SEQ1605 SEQ1606 SEQ1607 SEQ1608	TGATGATTTGGAACATGCGATTAAACTCTCTTTAGAAACTTTAAGTTTCCCAATTTTTG  AAAACCGGCTAATATGGGGTCATCAGTAGGTATTTCAAAAGCGACAGATGAATCCTCAC
SEQ1615 SEQ1616 SEQ1617 SEQ1601 SEQ1602 SEQ1603 SEQ1604 SEQ1605 SEQ1606 SEQ1607 SEQ1608 SEQ1609	TGATGATTTGGAACATGCGATTAAACTCTCTTTAGAAACTTTAAGTTTCCCAATTTTTG  AAAACCGGCTAATATGGGGTCATCAGTAGGTATTTCAAAAGCGACAGATGAATCCTCAC
SEQ1615 SEQ1616 SEQ1617 SEQ1601 SEQ1602 SEQ1603 SEQ1604 SEQ1605 SEQ1606 SEQ1607 SEQ1608 SEQ1609 SEQ1610	TGATGATTTGGAACATGCGATTAAACTCTCTTTAGAAACTTTAAGTTTCCCAATTTTTG  AAAACCGGCTAATATGGGGTCATCAGTAGGTATTTCAAAAGCGACAGATGAATCCTCAC
SEQ1615 SEQ1616 SEQ1617 SEQ1601 SEQ1602 SEQ1603 SEQ1604 SEQ1605 SEQ1606 SEQ1607 SEQ1608 SEQ1609 SEQ1610 SEQ1611 SEQ1611	TGATGATTTGGAACATGCGATTAAACTCTCTTTAGAAACTTTAAGTTTCCCAATTTTTG  AAAACCGGCTAATATGGGGTCATCAGTAGGTATTTCAAAAGCGACAGATGAATCCTCAC
SEQ1615 SEQ1616 SEQ1617 SEQ1601 SEQ1602 SEQ1603 SEQ1604 SEQ1605 SEQ1606 SEQ1607 SEQ1608 SEQ1609 SEQ1610 SEQ1611	TGATGATTTGGAACATGCGATTAAACTCTCTTTAGAAACTTTAAGTTTCCCAATTTTTG  AAAACCGGCTAATATGGGGTCATCAGTAGGTATTTCAAAAGCGACAGATGAATCCTCAC
SEQ1615 SEQ1616 SEQ1617 SEQ1601 SEQ1602 SEQ1603 SEQ1604 SEQ1605 SEQ1606 SEQ1607 SEQ1608 SEQ1609 SEQ1610 SEQ1611 SEQ1611 SEQ1612 SEQ1613	TGATGATTTGGAACATGCGATTAAACTCTCTTTAGAAACTTTAAGTTTCCCAATTTTTG  AAAACCGGCTAATATGGGGTCATCAGTAGGTATTTCAAAAGCGACAGATGAATCCTCAC
SEQ1615 SEQ1616 SEQ1617 SEQ1601 SEQ1602 SEQ1603 SEQ1604 SEQ1605 SEQ1606 SEQ1607 SEQ1608 SEQ1609 SEQ1610 SEQ1611 SEQ1611 SEQ1612 SEQ1613 SEQ1614	TGATGATTTGGAACATGCGATTAAACTCTCTTTAGAAACTTTAAGTTTCCCAATTTTTG  AAAACCGGCTAATATGGGGTCATCAGTAGGTATTTCAAAAGCGACAGATGAATCCTCAC
SEQ1615 SEQ1616 SEQ1617 SEQ1601 SEQ1602 SEQ1603 SEQ1604 SEQ1605 SEQ1606 SEQ1607 SEQ1608 SEQ1609 SEQ1610 SEQ1611 SEQ1611 SEQ1612 SEQ1613 SEQ1614 SEQ1615	TGATGATTTGGAACATGCGATTAAACTCTCTTTAGAAACTTTAAGTTTCCCAATTTTTG  AAAACCGGCTAATATGGGGTCATCAGTAGGTATTTCAAAAGCGACAGATGAATCCTCAC
SEQ1615 SEQ1616 SEQ1617 SEQ1601 SEQ1602 SEQ1603 SEQ1604 SEQ1605 SEQ1606 SEQ1607 SEQ1608 SEQ1609 SEQ1610 SEQ1611 SEQ1611 SEQ1612 SEQ1613 SEQ1614 SEQ1615 SEQ1616	TGATGATTTGGAACATGCGATTAAACTCTCTTTAGAAACTTTAAGTTTCCCAATTTTTG  AAAACCGGCTAATATGGGGTCATCAGTAGGTATTTCAAAAGCGACAGATGAATCCTCAC
SEQ1615 SEQ1616 SEQ1617 SEQ1601 SEQ1602 SEQ1603 SEQ1604 SEQ1605 SEQ1606 SEQ1607 SEQ1608 SEQ1609 SEQ1610 SEQ1611 SEQ1611 SEQ1612 SEQ1613 SEQ1614 SEQ1615 SEQ1616	TGATGATTTGGAACATGCGATTAAACTCTCTTTAGAAACTTTAAGTTTCCCAATTTTTG  AAAACCGGCTAATATGGGGTCATCAGTAGGTATTTCAAAAGCGACAGATGAATCCTCAC
SEQ1615 SEQ1616 SEQ1617 SEQ1601 SEQ1602 SEQ1603 SEQ1604 SEQ1605 SEQ1606 SEQ1607 SEQ1608 SEQ1609 SEQ1610 SEQ1611 SEQ1611 SEQ1612 SEQ1613 SEQ1614 SEQ1615 SEQ1615 SEQ1616	TGATGATTTGGAACATGCGATTAAACTCTCTTTAGAAACTTTAAGTTTCCCAATTTTTG  AAAACCGGCTAATATGGGGTCATCAGTAGGTATTTCAAAAGCGACAGATGAATCCTCAC
SEQ1615 SEQ1616 SEQ1617 SEQ1601 SEQ1602 SEQ1603 SEQ1604 SEQ1605 SEQ1606 SEQ1607 SEQ1608 SEQ1609 SEQ1610 SEQ1611 SEQ1611 SEQ1612 SEQ1613 SEQ1614 SEQ1615 SEQ1615 SEQ1616 SEQ1617	TGATGATTTGGAACATGCGATTAAACTCTCTTTAGAAACTTTAAGTTTCCCAATTTTTG  AAAACCGGCTAATATGGGGTCATCAGTAGGTATTTCAAAAGCGACAGATGAATCCTCAC
SEQ1615 SEQ1616 SEQ1617 SEQ1601 SEQ1602 SEQ1603 SEQ1604 SEQ1605 SEQ1606 SEQ1607 SEQ1608 SEQ1609 SEQ1610 SEQ1611 SEQ1611 SEQ1612 SEQ1613 SEQ1614 SEQ1615 SEQ1615 SEQ1616 SEQ1617 SEQ1601 SEQ1601	AAAACCGGCTAATATGGGGTCATCAGTAGTATCTCACAGAAAGCGACAGATGAAACCTCAC
SEQ1615 SEQ1616 SEQ1617 SEQ1601 SEQ1602 SEQ1603 SEQ1604 SEQ1605 SEQ1606 SEQ1607 SEQ1608 SEQ1609 SEQ1610 SEQ1611 SEQ1611 SEQ1612 SEQ1613 SEQ1614 SEQ1615 SEQ1615 SEQ1616 SEQ1617 SEQ1601 SEQ1601 SEQ1602 SEQ1603	AAAACCGGCTAATATGGGGTCATCAGTAGGTATTTCAAAAGCGACAGATGAATCCTCAC
SEQ1615 SEQ1616 SEQ1617 SEQ1601 SEQ1602 SEQ1603 SEQ1604 SEQ1605 SEQ1606 SEQ1607 SEQ1608 SEQ1609 SEQ1610 SEQ1611 SEQ1612 SEQ1613 SEQ1614 SEQ1615 SEQ1615 SEQ1616 SEQ1617 SEQ1601 SEQ1601 SEQ1602 SEQ1603 SEQ1604 SEQ1605	AAAACCGGCTAATATGGGGTCATCAGTAGGTATTTCAAAAGCGACAGATGAATCCTCAC
SEQ1615 SEQ1616 SEQ1601 SEQ1602 SEQ1603 SEQ1604 SEQ1605 SEQ1606 SEQ1607 SEQ1608 SEQ1609 SEQ1610 SEQ1611 SEQ1612 SEQ1613 SEQ1614 SEQ1615 SEQ1616 SEQ1617 SEQ1601 SEQ1601 SEQ1602 SEQ1603 SEQ1604 SEQ1605 SEQ1606	AAAACCGGCTAATATGGGGTCATCAGTAGTATTTCAAAAGCGACAGATGAATCCTCAC
SEQ1615 SEQ1616 SEQ1617 SEQ1601 SEQ1602 SEQ1603 SEQ1604 SEQ1605 SEQ1606 SEQ1607 SEQ1608 SEQ1609 SEQ1610 SEQ1611 SEQ1612 SEQ1613 SEQ1614 SEQ1615 SEQ1615 SEQ1616 SEQ1617 SEQ1601 SEQ1601 SEQ1602 SEQ1603 SEQ1604 SEQ1605	AAAACCGGCTAATATGGGGTCATCAGTAGGTATTTCAAAAGCGACAGATGAATCCTCAC
SEQ1615 SEQ1616 SEQ1601 SEQ1602 SEQ1603 SEQ1604 SEQ1605 SEQ1606 SEQ1607 SEQ1608 SEQ1610 SEQ1611 SEQ1612 SEQ1613 SEQ1614 SEQ1615 SEQ1616 SEQ1617 SEQ1601 SEQ1601 SEQ1602 SEQ1603 SEQ1604 SEQ1605 SEQ1606 SEQ1607	AAAACCGGCTAATATGGGGTCATCAGTAGTATTTCAAAAGCGACAGATGAATCCTCAC

# Table 16: Comparative Sequences relating t SAG0767 (D-alanine – D-alanine ligase)

	·
SEQ1610	ATGC-GTGCTATTAATTATGATAAATTTTTTTGTTAAAACTTATTTTATCACGCAAGTAG
SEQ1611	TCGCTCTGCAATTGACTTAGCTCTCAAGTATGATAGCCGTATTTTGATTGA
SEQ1612	ATGC-GTGCTATTAATTATGATAAATTTTTTTTTTAAAACTTATTTTATCACGCAAGTAG
SEQ1613	TCGCTCTGCAATTGACTTAGCTCTCAAGTATGATAGCCGTATTTTGATTGA
SEQ1614	ACCCCCCCCTABLIA ACCA ACCACCA ACCACCA ACCACCA ACCACCA ACCACC
SEQ1615	ATGCCGTGCTATTAATTATGATAAATTTTTTTTTTTAAAACTTATTTTATCACGCAAGTAG
	TCGCTCTGCAATTGACTTAGCTCTCAAGTATGATAGCCGTATTTTGATTGA
SEQ1616	ATGC-GTGCTATTAATTATGATAAATTTTTTGTTAAAACTTATTTTATCACGCAAGTAG
SEQ1617	TTGT-ATTCAGTTCGTTTAAGAAGATCTGTCCATCTTTCGTCAAAAAGAAATCACAGCG
SEQ1601	GACAGCTCGTGAAATTGAAGTAGGTATTTTAGGCAATAATGATGTTAAGACAACTTTTC
SEQ1602	GATAAACCACAAGCCCCGATTGCTTTAAAAGCTTTACTTGCATATTGACGCATTG
SEQ1603	GTCAATTTATTAAAACACAAGAATTTGATGAAATGCCATCTTCAGATGAAAAGTTA
SEQ1604	
SEQ1605	GTCAATTTATTAAAACACAAGAATTTGATGAAATGCCATCTTCAGATGAAAA
SEQ1606	GACAGCTCGTGAAATTGAAGTAGGTATTTTAGGCAATAATGATGTTAAGACAACTTTTC
SEQ1607	GACAGCTCGTGAAATTGAAGTAGGTATTTTAGGCAATAATGATGTTAAGACAACTTTTC
SEQ1608	GTCAATTTATTAAAACACAAGAATTTGATGAAAATGCCATCTTCAGATGAAAAGTTA
SEQ1609	GACAGCTCGTGAAATTGAAGTAGGTATTTTAGGCAATAATGATGTTAAGACAACTTTTC
SEQ1610	GTCAATTTATTAAAACACAAGAATTTGATGAAAATGCCATCTTCAGATGAAAAGTTA
SEQ1611	GACAGCTCGTGAAATTGAAGTAGGTATTTTAGGCAATAATGATGTTAAGACAACTTTTC
SEQ1612	GTCAATTTATTAAAACACAAGAATTTGATGATAATGCCATCTTCAGATGAAAAGTTA
SEQ1613	GACAGCTCGTGAAATTGAAGTATTTTTTTTTTTTTTTTT
SEQ1614	GTCAATTTATTAAAACACAAGAATTTGATGAAAATGCCATCTTCAGATGAAAAGTTA
SEQ1615	GLOCOTOTA A DETCA A CENTRAL CONTROL CO
SEQ1616	GACAGCTCGTGAAATTGAAGTAGGTATTTTTAGGCAATAATGATGTTAAGACAACTTTTC
SEQ1617	GTCAATTATTAAAACACAAGAATTTGATGAAATGCCATCTTCAGATGAAAAGTTA
BEQ1017	GATAAACCACAAGCCCCGATTGCTTTAAAAGCTTTACTTGCATATTGACGCATTG
SEQ1601	MCCCCA A CHIICHIII A A CA COMOCA PETERS
SEQ1602	TGGCGAAGTTGTTAAAGACGTCGATTTCTATGACTATGACGCCAAAT-ATATTGATA
SEQ1603	TTCCATAGTTGCTTCATCAACTTTAGCTGGAATATCCATAGTAATTTTATTATCA
SEQ1604	TGACAAACCAAACTGTTGATTTAGACAAAATGGTTCGTCCAAGTGATATCTATG
	CGTCGATTTCTATGACTATGACGCCAAAT-ATATTGATA
SEQ1605	
SEQ1606	TGGCGAAGTCGTTAAAGACGTCGATTTCTATGACTATGACGCCAAAT-ATATTGATA
SEQ1607	TGGCGAAGTCGTTAAAGACGTCGATTTCTATGACTATGACGCCAAAT-ATATTGATA
SEQ1608	TGACAAACCAAACTGTTGATTTAGACAAAATGGTTCGTCCAAGTGATATCTATG
SEQ1609	TGGCGAAGTCGTTAAAGACGTCGATTTCTATGACTATGACGCCAAAT-ATATTGATA
SEQ1610	TGACAAACCAAACTGTTGATTTAGACAAAATGGTTCGTCCAAGTGATATCTATG
SEQ1611	TGGCGAAGTCGTTAAAGACGTCGATTTCTATGACTATGACGCCAAAT-ATATTGATA
SEQ1612	TGACAAACCAAACTGTTGATTTAGACAAAATGGTTCGTCCAAGTGATATCTATG
SEQ1613	TGGCGAAGTTGTTAAAGACGTCGATTTCTATGACTATGACGCCAAAT-ATATTGATA
SEQ1614	TGACAAACCAAACTGTTGATTTAGACAAAATGGTTCGTCCAAGTGATATCTATG
SEQ1615	TGGCGAAGTCGTTAAAGACGTCGATTTCTATGACTATGACGCCAAAT-ATATTGATA
SEQ1616	TGACAAACCAAACTGTTGATTTAGACAAAATGGTTCGTCCAAGTGATATCTATG
SEQ1617	TTCCATAGATGCTTCATCAACTTTAGCTGGAATATCCATAGCAATTTTATTATCA
	The state of the s
SEQ1601	TAAAATTACTATGGATATTCCAGCTAAAGTTGATGAAGCAACTATGGAAGCAATGCG
SEQ1602	TATATTTGGCGTCATAGTCATAGAAATCGACGTCTTTAACGACTTCGCCAGGAAAAG
SEQ1603	TGATAATGCAATTGTTTTCCCCGTTTTACATGGACCAATGGGGGAAGATGGT
SEQ1604	TAAAATTACTATGGATATTCCAGCTAAAGTTGATGAAGCAACTATGGAAGCAATGCG
SEQ1605	
SEQ1606	TAAAATTACTAT — GGATATTCCA COMAAACTTCA TOO A COAT COAT COAT COAT COAT
SEQ1607	TAAAATTACTATGGATATTCCAGCTAAAGTTGATGAAGCAACTATGGAAGCAATGCG
SEQ1608	TAAAATTACTATGGATATTCCAGCTAAAGTTGATGAAGCAACTATGGAAGCAATGCG
SEQ1609	TGATAATGCAATTGTTTTCCCCGTTTTACATGGACCAATGGGGGAAGATGGT
SEQ1610	TAAAATTACTATGGATATTCCAGCTAAAGTTGATGAAGCAACTATGGAAGCAATGCG
	TGATAAT
SEQ1611	TAAAATTACTATGGATATTCCAGCTAAAGTTGATGAAGCAACTATGGAAGCAATGCG
SEQ1612	TGATAATGCAATTGTTTTCCCCGTTTTACATGGACCAATGGGGGAAGATGGT
SEQ1613	TAAAATTACTATGGATATTCCAGCTAAAGTTGATGAAGCAACTATGGAAGCAATGCG
SEQ1614	TGATAATGCAATTGTTTTCCCCGTTTTACATGGACCAATGGGGGAAGATGGT
SEQ1615	TAAAATTACTATGGATATTCCAGCTAAAGTTGATGAAGCAACTATGGAAGCAATGCG
SEQ1616	TGATAATGCAATTGTTTTCCCCGTTTTACATGGACCAATGGGGGAAGATGGT
SEQ1617	TATATTTGGCGTABLECMPARATIVESEQENCESRELA-TINGTSAGDALANI



SEQ1601	CAATATGCAAGTAAAGCTTTTAAAGCAATCGGGGCTTGTGGTTTATCACGCTGTGATTT
SEQ1602	TGTCTTAACATCATTATTGCCTAAAATACCTACTTCAATTTCACGAGCTGTCACGCC
SEQ1603	CTATCCAAGGATTTTTAGAAGTTTTAAGGATGCCTTATGTTGGGACTAATATTCT
SEQ1604	CAATATGCAAGTAAAGCTTTTAAAGCAATCGGGGCTTGTGGTTTATCACGCTGTGATTT
SEQ1605	
SEQ1606	CAATATGCAAGTAAAGCTTTTAAAGCAATCGGGGCTTGTGGTTTATCACGCTGTGATTT
SEQ1607	CAATATGCAAGTAAAGCTTTTAAAGCAATCGGGGCTTGTGGTTTATCACGCTGTGATTT
SEQ1608	CTATCCAAGGATTTTTAGAAGTTTTAAGGATGCCTTATGTTGGGACTAATATTCT
SEQ1609	CAATATGCAAGTAAAGCTTTTAAAGCAATCGGGGCTTGTGGTTTATCACGCTGTGATTT
SEQ1610	
SEQ1611	CAATATGCAAGTAAAGCTTTTAAAGCAATCGGGGCTTGTGGTTTATCACGCTGTGATTT
SEQ1612	CTATCCAAGGATTTTTAGAAGTTTTAAGGATGCCTTATGTTGGGACTAATATTCT
SEQ1613	CAATATGCAAGTAAAGCTTTTAAAGCAATCGGGGCTTGTGGTTTATCACGCTGTGATTT
SEQ1614	CTATCCAAGGATTTTTAGAAGTTTTAAGGATGCCTTATGTTGGGACTAATATTCT
SEQ1615	CAATATGCAAGTAAAGCTTTTAAAGCATCCGTTGTTGTTTTTTTT
SE01616	CAMIAIGCAAGIAAAGCTTTTAAAGCAATCGGGGCTTGTGGTTTATCACGCTGTGATTT
SEQ1617	CTATCCAAGGATTTTTAGAAGTTTTAAGGATGCCTTATGTTGGGACTAATATTCT
PEGIOIA	EDALANINELIGASE
CD01 C01	
SEQ1601	TTTTTGACGAAAGATGGACAAATCTTCTTAAACGAACTGAA-TACAATGCCCGGTTT
SEQ1602	TGTTCAATCAAAATACGGCTATCATACTTGAGAGCTAAGTCAATKSCAGAGCGAAGTGA
SEQ1603	TCTTCAAGCGTGGCTAT
SEQ1604	TTTTTGACGAAAGATGGACAAATCTTCTTAAACGAACTGAA-TACAATGCCC
SEQ1605	
SEQ1606	TTTTTGACGAAAGATGGACAAATCTTCTTAAACGAACTGAA-TACAATGCCCGGTTT
SEQ1607	TTTTTGACGAAAGATGGACAAATCTTCTTAAACGAACTGAA-TACAATGCCCGGTTT
SEQ1608	TCTTCAA
SEQ1609	TTTTTGACGAAAGAA-TGGACAAATCTTCTTAAACGAACTGAAATAC
SEQ1610	
SEQ1611	TTTTTGACGAAAGATGGACAAATCTTCTTAAACGAACTGAA-TACAATGCCCGGTTT
SEQ1612	TCTTCAAGCGTGGCTATGGATAAAATTACAACAAAACAA
SEQ1613	TTTTTGACGAAAGATGGACAAATCTTCTTAAACGAACTGAA-TACAATGCCCGGTTT
SEQ1614	TCTTCAAGCGTGGCTATGGATAAAATTACAACAAAACAA
SEQ1615	TTTTTGACGAAAGATGGACAAATCTTCTTAAACGAACTGAA-TACAATGCCCGGTTT
SEQ1616	TCTTCAAGCGTGGCTATGGATAAAATTACAACAAAACAA
SEQ1617	
SEQ1601	ACTCAGTGGTCAATGTATCCTCTGCTTTGGGAAAA-TATGGGGCTAACTTATAGTGATT
SEQ1602	GATTCATCTGTCGCTTTTGAAATACCTACTGATGACCCCATATTAGCCGGTTTTACAAA
SEQ1603	
SEQ1604	
SEQ1605	
SEQ1606	ACTCACTCA ATCTATCCACTCACTCACTCACTCACTCAC
SEQ1607	ACTCAGTGGTCAATGTATCCTCTGCTTTGGGAAAA-T
SEQ1608	ACTCAGTGGTCAATGTATCCCCTGCTTTGGGAAAAGTATGGGGCTAACCTT
SEQ1609	
SEQ1610	
SEQ1611	ACTCAGTGGTCAATGTATCCCCTGCTTTGGGAAAA-TATGGGGCTAACTTATAG
SEQ1612	
SEQ1613	ACTCAGTGGTCAATGTATCCTCTGCTTTGGGAAAA-TATGGGGCTAACTT
SEQ1614	CCTCAGG
SEQ1615	ACTCAGTGGTCAATGTATCCCCTGCTTTGGGAAAA-TATGGGGCTAACTTATAGTGA
SEQ1616	
SEQ1617	

# Table 16: Comparative Sequences relating to SAG0767 (D-alanine – D-alanine ligase)

SEQ1601	GATTG
SEQ1602	ATTGGGAAACTTAAAGTTTCTAAAGAGAGTTTAATCGCATGTTCCAAATCATCACCCTC
SEQ1603	
SEQ1604	
SEQ1605	
SEQ1606	
SEQ1607	
SEQ1608	***
SEQ1609	
SEQ1610	
SEQ1611	
SEQ1612	
SEQ1613	
SEQ1614	▼ · · · · · · · · · · · · · · · · · · ·
SEQ1615	***************************************
SEQ1616	
SEQ1617	
SEQ1017	
SEQ1601	
SEQ1602	7.7.7.7.7.0000000000000000000000000000
	AAATAAGTTTGATATGCAACCTGAGGTACACCTACTGTTGCAAGGACTTGTTTTGTTGT
SEQ1603	
SEQ1604	
SEQ1605	
SEQ1606	
SEQ1607	,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,
SEQ1608	
SEQ1609	
SEQ1610	#===+=================================
SEQ1611	
SEQ1612	
SEQ1613	
SEQ1614	
SEQ1615	
SEQ1616	~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~
SEQ1617	
SEQ1601	**************************************
SEQ1602	ATTTTATCCATAGCCACGCTTGAAGATAGAATATTAGTCCCAACATAAGGCATCCTTAA
SEQ1603	
SEQ1604	, 
SEQ1605	
SEQ1606	
SEQ1607	
SEQ1608	
SEQ1609	*
SEQ1610	
SEQ1611	
SEQ1612	
SEQ1613	***************************************
SEQ1614	
SEQ1615	
SEQ1616	*
SEQ1617	

## Sable 16: Comparative Sequences relating to SAG0767 (D-alanine – D-alanine ligase)

SEQ1601	
SEQ1602	ACTTCTAAAAATCCTTGGATAGAACCATCTTCCCCCATTGGTCCATGTAAAACGGGGAA
SEQ1603	
SEQ1604	~
SEQ1605	
SEQ1606	
SEQ1607	
SEQ1608	
SEQ1609	
SEQ1610	
SEQ1611	
SEQ1612	
SEQ1613	
SEQ1614	
SEQ1615	
SEQ1616	
SEQ1617	
PEGICIA	
SEQ1601	
SEQ1601 SEQ1602	ACAATTGCATTATCATCATAGATATCACTTGGACGAACCATTTTGTCTAAATCAACAGT
	ACAATTGCATTATCATCATAGATATCACTTGGACGAACCATTTTGTCTAAATCACAGT
SEQ1603	
SEQ1604	
SEQ1605	
SEQ1606	
SEQ1607	
SEQ1608	
SEQ1609	
SEQ1610	
SEQ1611	
SEQ1612	
SEQ1613	
SEQ1614	
SEQ1615	
SEQ1616	
SEQ1617	
SEQ1601	
SEQ1602	TGGTTTGTCATTAACTTTTCATCTGAAGATGGCATTTCATCAAATTCTTGTGTTTTAAT
SEQ1603	
SEQ1603	#======;
SEQ1605	
SEQ1605	
SEQ1606 SEQ1607	
SEQ1608	
SEQ1609	
SEQ1610	
SEQ1611	
SEQ1612	
SEQ1613	
SEQ1614	
SEQ1615	
SEQ1616	
SEQ1617	
<del>-</del>	

# Table 16: Comparative Sequences relating to SAG0767 (D-alanine – D-alanine ligase)

SEQ1601	
SEQ1602	AATTGACCTACTTGCGTG
SEQ1603	***********
SEQ1604	
SEQ1605	
SEQ1606	
SEQ1607	
SEQ1608	
SEQ1609	*=
SEQ1610	
SEQ1611	
SEQ1612	
SEQ1613	
SEQ1614	
SEQ1615	
SEQ1616	
SEO1617	



## Table 17: C mparative Sequences relating to SAG1086 (xanthine ph phorib syltransferase)

SEQ ID NO. 1701: SAG1086 FROM THE1169NT1 GBS NONTYPEABLE STRAIN
TTTANAGGTTGATTCCTTTTTGACTCATCAGGTAGATTTTGAGTTAATGCAGGAAATAGGTAAAGTTTTTGCTGATAAATATAAAGAAG
CCGGCATTACGAAGGTTGTTACGATTGAAGCATCTGGAATTGCGCCAGCAGTGTACGCAGCTCAAGCATTGGGCGTACCAATCATATTT
GCTAAAAAGGCTAAGAACATTACTATGACTGAAGGTATCTTAACTGCTGAAGTGTATTCTTTTACAAAGCAAGGTTCAAGTTTC
TATTGTGAGTCGCTTTTTATCTAACGATGATACTGTACTCATCATTGATGACTTTTTAGCAAACGGTCAAGCGGCTAAAGGATTACTTG
AAAATTATTGGTCAAGCTGGAGCTAAGGTTGCTGGTATCGGAATCGTTATTGAAAAAATCTTTCCAAGATGGGCGTGATTTGTTAGAAAAA
ACAGGTGTTCCAGT

SEQ ID NO. 1702: SAG0767 FROM THE 18RS21 GBS TYPE II STRAIN
TTTAGGTGAGAACATTTTAAAGGTTGATTCTTTTTTGACTCATCAGGTAGATTTTGAGTTAATGCAGGAAATAGGTAAAGTTTTTGCTG
ATAAATATAAAGAAGCCGGCATTACGAAGGTTGTTACGATTGAAGCATCTGGAATTGCACCAGCAGTGTACGCAGCTCAAGCATTGGGC
GKACCAATGATATTTGCTAAAAAAGCTAAGAACATTACTATGACTGAAGGTATCTTAACTGCTGAAGTGTATTCTTTTACAAAACGAAGCT
TACGAGTCAAGTTTCTATTGTGAGTCGCTTTTTATCTAACGATGATACTGTACTCATCATTGATGACTTTTTAGCAAACGGTCAAGCGG
CTAAAGGATTACTTGAAATTATTGGTCAAGCTGGAGCTAAGGTTGCTGGTATCGGAATCGTTATTGAAAATACTTTCCAAGATGGGCGT
GATTTGTTAGAAAAAAACA

SEQ ID NO. 1703: SAG0767 FROM THE H36b1 GBS TYPE Ib STRAIN
AAGAACGTATTCTTAAAGATGGTGATGTTTTAGGTGAĞAACATTTTAAAAGTTGATTCTTTTTTGACTCATCAGGTAGATTTTGAGTTA
ATGCAGGAAATAGGTAAAGTTTTTGCTGATAAATATAAAGAAGCCGGCATTACGAAGGTTGTTACAATTGAAGCATCTGGAATTGCGCC
AGCAGTGTACGCAGCTCAAGCATTGGGCGTACCAATGATATTTGCTAAAAAAGCTAAGAACATTACTATGACTGAAGTATCTTAACTG
CTGAAGTGTATTCTTTTTACAAAGCAAGCTAAGGATCAAGTTACTTGAAATTATTGGTCAAGCTGAGCTTAACGATGATCTTTACTATCATCATT
TATTGAAAAACGTCAAGATGGGCCTAAAGGATTACTTGAAATTATTGGTCAAGCTGGAGCTAAGGTTGCTGGTATCGGAATCYT
TATTGAAAAAATCTTTCCAAGATGGGCGTGATT

SEQ ID NO. 1704: SAG0767 FROM THE M732 GBS TYPE III STRAIN
ATTCTTTTTTGACTATCAGGTAAATTTTGAGTTAATGCAGGAAATAGGTAAAGTTTTTGCTGATAAATATAAAGAAGCCGGCATTACGA
AGGTTGTTACAATTGAAGCATCTGGAATTGCGCCAGCAGTGTACGCAGCTCAAGCATTGGGCGTACCAATGATATTTGCTAAAAAAGCT
AAGAACATTACTATGACTGAAGGTATCTTAACTGCTGAAGTGTATTCTTTTACAAAGCAAGTTACGAGTCAAGTTTCTATTGTAGTCG
CTTTTTATCTAACGATGATACTGTACTCATCATTGATGACTTTTTAGCAAACGGTCAAGCGGCTAAAGGATTACTTGAAATTATTGGTC
AAGCTGAAGCTAAGGTTGCTGGTATCGGAATCGTTATTGAAAAAATCTTTCCAAGATGGGCGTGATTTGTTAGAAAAAAACAGGTGTTCCG
GTTACTTCTCTTGCTCGT

SEQ ID NO. 1705: SAG0767 FROM THE M781 GBS TYPE III STRAIN REVERSE COMPLEMENT GAACGTATTCTTAAAGATGGTGATGTTTTTAGGTGAGAACATTTTAAAAGTTGATTCTTTTTTTGACTCATCAGGTAAATTTTGAGTTAAT GCAGGAAATAGGTAAAGTTTTTGCTGATAAATATAAAGAAGCCGGCATTACGAAGGTTGTTACAATTGAAGCATCTGGAATTGCGCCAG CAGTGTACGCAGCCTCAAGCATTGGGCGTACCAATGATATTTGCTAAAAAAGCTAAGAACATTACTATGACTGAAGGTATTTAACTGCT GAAGTGTATTCTTTTACAAAGCAAGCTACCATCATTGA TGACTTTTTAACAAACGGTCAAGC

SEQ ID NO. 1706: SAG0767 FROM THE 090 GBS TYPE IA STRAIN REVERSE COMPLEMENT ACATTTTAAAGGTTGATTCTTTTTTGACTCATCAGGTAGATTTTGAGTTAATGCAGGAAATAGGTAAAGTTTTTGCTGATAAATATAAA GAAGCCGGCATTACGAAGGTTGTTACGATTGAAGCATCTGGAATTGCACCAGCAGTGTACGCAGCTCAAGCATTGGGCGTACCAATGAT ATTTGCTAAAAAAGCTAAGAACATTACTATGACTGAAGGTATCTTAACTGCTGAAGTGTATTCTTTTACAAAGCAAGTTACGAGTCAAGTTTCTATTGTAGATCAGCTTTTTATCTAACGATGATACTGATCATCATTGATGACTTTTTAGCAAACMGTCYAGCGGCTAAAGGATTA CTTGAAAATTTTTGGTCAAGGTGGAGCTGATGTGTTAGGAATCGTTATTGAAAAATCTTTCCAAGATGGGCGTGATTTGTTAGAAAA

SEQ ID NO. 1709: SAG0767 FROM THE CJB110 GBS NONTYPEABLE STRAIN REVERSE COMPLEMENT

GCTGATAAATATAAAGAAGCCGGCATTACGAAGGTTGTTACAATTGAAGCATCTGGAATTGCGCCAGCAGTGTACGCAGCTCAAGCATT
GGGCGTACCAATGATATTTGCTAAAAAAGCTAAGAACATTACTATGACTGAAGGTATCTTAACTGCTGAAGTGTATTCTTTTACAAAAGC
AAGTTACGAGTCAAGTTTCTATTGTGAGTCGCTTTTTATCTAACGATGATACTGTACTCATCATTGATGACTTTTTAGCAAACGGTCAA
GCGGCTAAAGGATTACTTGAAATTTATTGGTCAAGCTGGAGCTAAGGTTGCTGGTATCGGAATCGTTATTGAAAAAACTTTTCCAAGATG
GGCGTGATTTGTTAGAAAAAACAGGTGTTCCAGT



### SEQ ID NO. 1710: SAG0767 FROM THE 2603 V/R GBS TYPE V STRAIN

# SEQ ID NO. 1711: SAG0767 FROM THE JM9130013 GBS TYPE VIII STRAIN REVERSE COMPLEMENT

,	
SEQ1701	TTTAAAGGTTGATTCCT
SEQ1702	TTTAGGTGAGAACATTTTAAGGTTGATTCTT
SEQ1703	AGAACGTATTCTTAAAGATGGTGATGTTTTAGGTGAGAACATTTTAAAAGTTGATTCTT
SEQ1704	
SEQ1705	-GAACGTATTCTTAAAGATGGTGATGTTTTAGGTGAGAACATTTTAAAAAACATGCTTT
SEQ1706	ACATTTTAAAGGTTGATTCTT
SEQ1707	ACGTATTCTTAAAGATGGTGATGTTTTAGGTGAGAACATTTTAAAAGTTGATTCTT
SEQ1708	TTTAAAAGTTGATTCTT
SEQ1709	TITAAAGITGATICTI
SEQ1710	AACGTATTCTTAAAGATGGTGATGTTTTAGGTGAGAACATTTTAAAAGTTGATTCTT
SEQ1711	THE CONTROL OF THE CO
SEQ1701	TTTGACTCATCAGGTAGATTTTGAGTTAATGCAGGAAATAGGTAAAGTTTTTGCTGATA
SEQ1702	TTTGACTCATCAGGTAGATTTTGAGTTAATGCAGGAAATAGGTAAAGTTTTTGCTGATA TTTGACTCATCAGGTAGATTTTTGAGTTAATGCAGGAAATAGGTAAAGTTTTTTGCTGATA
SEQ1703	TTTGACTCATCAGGTAGATTTTGAGTTAATGCAGGAAATAGGTAAAGTTTTTGCTGATA TTTGACTCATCAGGTAGATTTTTGAGTTAATGCAGGAAATAGGTAAAGTTTTTTGCTGATA
SEQ1704	TTTTGACTATCAGGTAGATTTTGAGTTAATGCAGGAAATAGGTAAAGTTTTTGCTGATA TTTTGACTATCAGGTAAATTTTTGAGTTAATGCAGGAAATAGGTAAAGTTTTTTGCTGATA
SEQ1705	TTTGACTATCAGGTAAATTTTGAGTTAATGCAGGAAATAGGTAAAGTTTTTGCTGATA TTTGACTCATCAGGTAAATTTTTGAGTTAATGCAGGAAATAGGTAAAGTTTTTTGCTGATA
SEQ1706	TITICACICATCA CON COMPANY DE CONTROL C
SEQ1707	TTTGACTCATCAGGTAGATTTTGAGTTAATGCAGGAAATAGGTAAAGTTTTTGCTGATA
SEQ1707	TTTGACTCATCAGGTAGATTTTGAGTTAATGCAGGAAATAGGTAAAGTTTTTGCTGATA
SEQ1709	TTTGACTCATCAGGTAAATTTTGAGTTAATGCAGGAAATAGGTAAAGTTTTTGCTGATA
SEQ1709 SEQ1710	The state of the s
SEQ1711	TTTGACTCATCAGGTAGATTTTGAGTTAATGCAGGAAATAGGTAAAGTTTTTGCTGATA
35Q1/11	
SEQ1701	A TATA A ACA D COCCOCA BUILD COS P COMPONENT COS TRACA DO COSTA COS TRACA DO COSTA COS TRACA DO COSTA COS TRACA DO COSTA
SEQ1702	ATATAAAGAAGCCGGCATTACGAAGGTTGTTACGATTGAAGCATCTGGAATTGCGCCAG
SEQ1702	ATATAAAGAAGCCGGCATTACGAAGGTTGTTACGATTGAAGCATCTGGAATTGCACCAG
SEQ1703	ATATAAAGAAGCCGGCATTACGAAGGTTGTTACAATTGAAGCATCTGGAATTGCGCCAG
SEQ1704 SEQ1705	ATATAAAGAAGCCGGCATTACGAAGGTTGTTACAATTGAAGCATCTGGAATTGCGCCAG
SEQ1705	ATATAAAGAAGCCGGCATTACGAAGGTTGTTACAATTGAAGCATCTGGAATTGCGCCAG
-	ATATAAAGAAGCCGGCATTACGAAGGTTGTTACGATTGAAGCATCTGGAATTGCACCAG
SEQ1707	ATATAAAGAAGCCGGCATTACGAAGGTTGTTACAATTGAAGCATCTGGAATTGCGCCAG
SEQ1708	ATATAAAGAAGCCGGCATTACGAAGGTTGTTACAATTGAAGCATCTGGAATTGCGCCAG
SEQ1709	ATATAAAGAAGCCGGCATTACGAAGGTTGTTACAATTGAAGCATCTGGAATTGCGCCAG
SEQ1710	ATATAAAGAAGCCGGCATTACGAAGGTTGTTACAATTGAAGCATCTGGAATTGCGCCAG
SEQ1711	ACGAAGGTTGTTACAATTGAAGCATCTGGAATTGCGCCAG
0001501	
SEQ1701	CAGTGTACGCAGCTCAAGCATTGGGGCGTACCAATGATATTTGCTAAAAAGGCTAAGAACA
SEQ1702	CAGTGTACGCAGCTCAAGCATTGGGCGKACCAATGATATTTGCTAAAAAAGCTAAGAACA
SEQ1703	CAGTGTACGCAGCTCAAGCATTGGGCGTACCAATGATATTTGCTAAAAAAGCTAAGAACA
SEQ1704	CAGTGTACGCAGCTCAAGCATTGGGCGTACCAATGATATTTGCTAAAAAAGCTAAGAACA
SEQ1705	CAGTGTACGCAGCTCAAGCATTGGGCGTACCAATGATATTTGCTAAAAAAGCTAAGAACA
SEQ1706	CAGTGTACGCAGCTCAAGCATTGGGCGTACCAATGATATTTGCTAAAAAAGCTAAGAACA
SEQ1707	CAGTGTACGCAGCTCAAGCATTGGGCGTACCAATGATATTTGCTAAAAAAGCTAAGAACA
SEQ1708	CAGTGTACGCAGCTCAAGCATTGGGCGTACCAATGATATTTGCTAAAAAAGCTAAGAACA
SEQ1709	CAGTGTACGCAGCTCAAGCATTGGGCGTACCAATGATATTTGCTAAAAAAGCTAAGAACA
SEQ1710	CAGTGTACGCAGCTCAAGCATTGGGCGTACCAATGATATTTGCTAAAAAAGCTAAGAACA
SEQ1711	CAGTGTACGCAGCTCAAGCATTGGGCGTACCAATGATATTTGCTAAAAAAGCTAAGAACA
	THE

# Table 17: Comparative Sequences relating t SAG1086 (xanthine phophoribosyltransferase)

	·
SEQ1701	TTACTATGACTGAAGGTATCTTAACTGCTGAAGTGTATTCTTTTACAAAGCAAGWTACGA
SEQ1702	TTACTATGACTGAAGGTATCTTAACTGCTGAAGTGTATTCTTTTACAAAGCAAGTTACGA
	THE COLORS OF TH
SEQ1703	TTACTATGACTGAAGGTATCTTAACTGCTGAAGTGTATTCTTTTACAAAGCAAGTTACGA
SEQ1704	TTACTATGACTGAAGGTATCTTAACTGCTGAAGTGTATTCTTTTACAAAGCAAGTTACGA
SEQ1705	TTACTATGACTGAAGGTATRTTAACTGCTGAAGTGTATTCTTTTACAAAGCAAGTTACGA
SEQ1706	TTACTATGACTGAAGGTATCTTAACTGCTGAAGTGTATTCTTTTACAAAGCAAGTTACGA
	TIACIAIGACIGAAGGIATCI TAACIGCIGAAGIGTAT TCTTTTACAAAGCAAGTTACGA
SEQ1707	TTACTATGACTGAAGGTATCTTAACTGCTGAAGTGTATTCTTTTACAAAGCAAGTTACGA
SEQ1708	TTACTATGACTGAAGGTATCTTAACTGCTGAAGTGTATTCTTTTACAAAGCAAGTTACGA
SEQ1709	TTACTATGACTGAAGGTATCTTAACTGCTGAAGTGTATTCTTTTACAAAGCAAGTTACGA
SEQ1710	TTACTATGACTGAAGGTATCTTAACTGCTGAAGTGTATTCTTTTACAAAGCAAGTTACGA
	I TACIATGACTGAAGGTATCTTAACTGCTGAAGTGTATTCTTTTACAAAGCAAGTTACGA
SEQ1711	TTACTATGACTGAAGGTATCTTAACTGCTGAAGTGTATTCTTTTACAAAGCAAGTTACGA
SEQ1701	GTCAAGTTTCTATTGTGAGTCGCTTTTTATCTAACGATGATACTGTACTCATCATTGATG
SEQ1702	CMCA A CHIMICAD AMICANCA CHICAGAMATA HOMA A CAN HOLA CAN CHICA TO CAN CHILA CAN CHICAGAMATA CAN HOLA CAN CHICAGAMATA CAN HOLA CAN CHICAGAMATA
	GTCAAGTTTCTATTGTGAGTCGCTTTTTATCTAACGATGATACTGTACTCATCATTGATG
SEQ1703	GTCAAGTTTCTATTGTGAGTCGCTTTTTATCTAACGATGATACTGTACTCATCATTGATG
SEQ1704	GTCAAGTTTCTATTGTGAGTCGCTTTTTATCTAACGATGATACTGTACTCATCATTGATG
SEQ1705	GTCAAGTTTCTATTGTGAGTCGCTTTTTATCTAACGATGATACTGTACTCATCATTGATG
SEQ1706	
	GTCAAGTTTCTATTGTGAGTCGCTTTTTATCTAACGATGATACTGTACTCATCATTGATG
SEQ1707	GTCAAGTTTCTATTGTGAGTCGCTTTTTATCTAACGATGATACTGTACTCATCATTGATG
SEQ1708	GTCAAGTTTCTATTGTGAGTCGCTTTTTATCTAACGATGATACTGTACTCATCATTGATG
SEQ1709	GTCAAGTTTCTATTGTGAGTCGCTTTTTATCTAACGATGATACTGTACTCATCATTGATG
SEQ1710	GTCAAGTTTCTATTGTGAGTCGCTTTTTATCTAACGATGATACTGTACTCATCATTGATG
SEQ1711	GTCAAGTTTCTATTGTGAGTCGCTTTTTATCTAACGATGATACTGTACTCATCATTGATG
	,
SEQ1701	ACTTTTTAGCAAACGGTCAAGCGGCTAAAGGATTACTTGAAATT-ATTGGTCAAGCTGGA
SEQ1702	ACTTTTTAGCAAACGGTCAAGCGGCTAAAGGATTACTTGAAATT-ATTGGTCAAGCTGGA
SEQ1703	ACTTTTTAGCAAACGGTCAAGCGGCTAAAGGATTACTTGAAATT-ATTGGTCAAGCTGGA
SEQ1704	ACTTTTTAGCAAACGGTCAAGCGGCTAAAGGATTACTTGAAATT-ATTGGTCAAGCTGAA
SEQ1705	ACTTTTTAACAAACGGTCAAGC
SEQ1706	ACTTTTTAGCAAACMGTCYAGCGGCTAAAGGATTACTTGAAATT-ATTGGTCAAGCTGGA
SEQ1707	ACTTTTTAGCAAACGGKCAAGCGGSTAAAGGATTACTTGAAATT-ATTGGTCAAGCTGGA
SEQ1708	ACTTTTTAGCAAACGGTCAAGCGGCTAAAGGATTACTTGAAATT-ATTGGTCAAGCTGAA
SEQ1709	ACTTTTTAGCAAACGGTCAAGCGGCTAAAGGATTACTTGAAATTTATTGGTCAAGCTGGA
SEQ1710	ACTTTTTAGCAAACGGTCAAGCGGCTAAAGGATTACTTGAAATT-ATTGGTCAAGCTGGA
SEQ1711	ACTTTTTAGCAAACGGTCAAGCGGCTAAAGGATTACTTGAAATT-ATTGGTCAAGCTGGA
SEQ1701	CTAAGGTTGCTGGTATCGGAATCGTTATTGAAAAATCTTTCCAAGATGGGCGTGATTTG
SEQ1702	
	CTAAGGTTGCTGGTATCGGAATCGTTATTGAAAAATCTTTCCAAGATGGGCGTGATTTG
SEQ1703	CTAAGGTTGCTGGTATCGGAATCYTTATTGAAAAATCTTTCCAAGATGGGCGTGATT
SEQ1704	CTAAGGTTGCTGGTATCGGAATCGTTATTGAAAAATCTTTCCAAGATGGGCGTGATTTG
SEQ1705	
SE01706	CTAAGGTTGCTGGTATCGGAATCGTTATTGAAAAATCTTTCCAAGATGGGCGTGATTTG
SEQ1707	CTA
SEQ1708	CTAAGGTTGCTGGTATCGGAATCGTTATTGAAAAATCTTTCCAAGATGGGCGTGATTTG
SE01709	CTAAGGTTGCTGGTATCGGAATCGTTATTGAAAAATCTTTCCAAGATGGGCGTGATTTG
SEQ1710	OTA A COMMOCOM MOCOTA MOCOTANIA MODERNA DE LA PROMINCIA DE MOCOTA DE MOCOTA MOC
	CTAAGGTTGCTGGTATCGGAATCGTTATTGAAAAATCTTTCCAAGATGGGCGTGATTTG
SEQ1711	CTAAGGTTGCTGGTATCGGATABCMARATVSTNCSRATNGTSAGXANTHN
SEQ1701	TAGAAAAAACAGGTGTTCCAGT
SEQ1702	TAGAAAAACA
_	ANDREMORAL TO THE TOTAL TO THE TOTAL
SEQ1703	
SEQ1704	TAGAAAAACAGGTGTTCCGGTTACTTCTCTTGCTCGT
SEQ1705	
SEQ1706	TAGAAAA
SEQ1707	######################################
SEQ1708	TAGAAAAACAGGTGTTCCGGTTAC
SEQ1709	TAGAAAAACAGGTGTTCCAGT
SEQ1710	TAGAAAAACAGGTGTTCCAG
SEQ1711	HRBSYTRANSRAS

## Table 18: Comparativ Sequences relating to SAG1600 (glutamate racemase)

SEQ ID NO. 1802: SAG1600 FROM THE M732 GBS TYPE III STRAIN REVERSE COMPLEMENT
AAATGTTCCGTCAACTTCCAGAAGAGAGAGTAATCTTCATTGGAGATCAGGCTAGAGCTCCGTATGGTCCTAGACCTGCTCAACAGATT
AGAGAGTTTACCTGGCAGATGGTTAACTTCTTATTGACTAAAAATGTTAAGATGATTGTTATAGCTTGTAATACAGCAACTGCAGTTGC
CTGGCAAGAAATTAAAGAAAAACTAGACATCCCTGTTTTAGGCGTTATTTTACCAGAGCTTAGCGCAGCTATCAAATCAACTAATTTAG
GGAAAGTTGGTATATAGGTACTCCCATGATTGTTAAATCAGATGCTTATCGTCAAAAAAATTCAAGCTTTGTCCCAAATACTGCTGTG
GTATCCCTTGCTTGCTCGAAATTTGTTCCAATTTTTAGGTTGCACGCATTATCCCCTATTAGCCAAAAAGGTGGTTTATAGAACGTT
GTCCCCATTAGTTGGTAAATTAGATACTTTAATTTTAGGTTGCACGCATTATCCCCTATTACGTCCCATCATTCAAAATGTTATGGGGG
CTGAGGTTAAATTAATTGATAGTGGCGCAGAAACCGTTCGTGATATTCTGTTTTATTGAACTATTTTGAGATAAACCATAATTGGCAA
AATAAACACGGTGGTCATCACTTTTACACAACCGCCAGCCCAAAAGGTTTTAAAGAAA

SEQ ID NO. 1805: SAG1600 FROM THE COHI GBS TYPE IA STRAIN
TTCCGTCAACTTCCAAAATATGAAGTAATCTTCATTGGAGATCAGGCTAGAGCTCCGTATGGTCCTAGACCTGCTCAACAGATTAGAGA
GTTTACCTGGCAGATGGTTAACTTCTTATTGACTAAAAATGTTAAGATGATTGTTATAGCTTGTAATACAGCAACTGCAGTTGCCTGGC
AAGAAATTAAAGAAAAACTAGACATCCCTGTTTTAGGCGTTATTTTACCAGGAGCTAGCGCAGCTATCAAATCAACTAATTTAGGGAAA
GTTGGTATTATAGGTACTCCCATGACTGTTAAATCAGATGCTTATCGTCAAAAAATTCAAGCTTTGTCTCCAAAATACTGCTGTGGTATC
CCTTGCTTGTCCCGAAAT

SEQ ID NO. 1806: SAG1600 FROM THE CJB110 GBS NONTYPEABLE STRAIN
GTAATCTTCATTGGAGATCAGGCTAGAGCTCCGTATGGTCCTAGACCTGCTCAACAGATTAGAGAGTTTACCTGGCAGATGGTTAATTT
CTTATTGACTAAAAATGTTAAGATGATTATAGCTTGTAATACAGCAACTGCAGTTGCCTGGCAAGAAATTAAAGAAAAACTAGACA
TAC

SEQ ID NO. 1808: SAG1600 FROM THE 1169NT1 GBS TYPE V STRAIN
GTAATCTTCATTGGGGATCAGGCTAGAGCTCCGTATGGTCCTAGACCTGCTCAACAGATTAGAGAGTTTACCTGGCAGATGGTTAATTT
CTTATTGACTAAAAATGTTAAGATGATTGTTATAGCTTGTAATACAGCAACTGCAGTT

SEQ ID NO. 1809: SAG1600 FROM THE 18RS21 GBS TYPE II STRAIN
GARATGTTCCGTCAACTTCCAGAAGAGGAAGTAATCTTCATTGGAGATCAGGCTAGAGCTCCGTATGGTCCTAGACCTGCTCAACAGAT
TAGAGAGTTTACCTGGCAGATGGTTAACTTCTTATTGACTAAAAATGTTAAGATGATTGTTATAGCTTGTAATACAGCAACTGCAGTTG
CCTGGCAAGAAATTAAAGAAAAACTAGACATCCCTGTTTTAGGCGTTATTTTACCAGGAGCTAGCGCAGCTATCAAATCAACTAATTTA
GGGAAAGTTGGTATTATAGGTACTCCCATGACTGTTAAATCAGATGCTTATCGTCAAAAAATTCAAGC



### Table 18: Comparative Sequences relating to SAG1600 (glutamate racemase)

SEQ ID NO. 1810: SAG1600 FROM THE 18RS21 TYPE II STRAIN

SEQ ID NO. 1811: SAG1600 FROM THE 2603 V/R GBS TYPE V STRAIN

SEQ ID NO. 1812: SAG1600 FROM THE M781 GBS TYPE III STRAIN

GGCGGTTGTGTAAAAGTGATGACCACCGTGTTTATTTTGCCAATTATGGTTTATCTCAAAAATAGTTCAATAAAACAGAAATATCACGAA CGGTTTCTGCGCCACTATCAATTAATTTAACCTCAGCCCCCATAACATTTTGAATGATGGGGACGTAATAGGGGATAATGCGTGCAACCT AAAATTAAAGTATCTAATTTACCAACTAATGGGGACAACGTTTCATAAACCACCTTTTTGGCTAAACTAGAAGA

SEQ ID NO. 1813: SAG1600 FROM THE M 781 GBS TYPE III STRAIN

AATCTTCATTGGAGATCAGGCTAGAGCTCCGTATGGTCCTAGACCTGCTCAACAGATTAGAGAGTTTACCTGGCAGATGGTTAACTTCT
TATTGACTAAAAATGTTAAGATGATTGTTATAGCTTGTAATACAGCAACTGC

SEQ ID NO. 1814: SAG1600 FROM THE JM9130013 GS TYPE VIII STRAIN

TGGGCTGGCGGTTGTGTAAAAGTGATGACCACCGTGTTTATTTTGCCAATTATGGTTTATCTCAAAATAGTTCAATAAAACAGAAATAT CACGAACGGTTTCTGCGCCACTATCAATTAATTTAACCTCAGCCCCCCATAACATTTTGAATGATGGGACGTAATAAGGGATAATGCGTG CAACCTAAAATTAAAGTATCTAATTTACCAACTAATGGGGACAACGTTTCATAAACCACCTTTTTGGCTAAACTAGAAGACATCTGATT TGATTCCACAATTGGAACAAATTTCGGACAAGCAAGGGATACCACACGCAGTATTTGGAGACAAAGCTTGAATTTTTTGACGATAAGCAT CTGATTTAACAGTCATGGGAGTACCTATAATACCAACTTTCCCTGAA

SEQ1801	AATCTTCATTGGAGATCAGGCTAGAGCT
SEQ1802	AAATGTTCCGTCAACTTCCAGAAGAGGAAGTAATCTTCATTGGAGATCAGGCTAGAGCT
SEQ1803	AATCTTCATTGGAGACCAGGCTAGAGCT
SEQ1804	GCGGTTGTGTAAAAG-T
SEQ1805	ttccgtcaacttccaaaatatgaagtaatcttcattggagatcaggctagagct
SEQ1806	gtaatcttcattggagatcaggctagagct
SEQ1807	CTTTTGGGCTGGCGGTTGTGTAAAAT-T
SEQ1808	gtaatcttcattggggatcaggctagagct
SEQ1809	AAATGTTCCGTCAACTTCCAGAAGAGGAAGTAATCTTCATTGGAGATCAGGCTAGAGCT
SEQ1810	ATTTCTTTAAAACCTTTTGGGCTGGCGGTTGTGTAATAT-T
SEQ1811	ATTTCTTTAAAACCTTTTGGGCTGGCGTTGTGTAATAAGT
SEQ1812	GGCGGTTGTGTAAAAG-T
SEQ1813	AATCTTCATTGGAGATCAGGCTAGAGCT
SEQ1814	TGGGCTGGCGGTTGTGTAAAAG-T
SEQ1801	CGTATGGTC-CTAGACCTGCTCAACAGATTAGAGAGTTTACCTGGCAGATGGTTAATTT
SEQ1802	CGTATGGTC-CTAGACCTGCTCAACAGATTAGAGAGTTTACCTGGCAGATGGTTAACTT
SEQ1803	CGTATGGTC-CTAGACCTGCTCAACAGATTAGAGAGTT-ACCTGGCAGATGGTTAATTT
SEQ1804	GATGACCACCGTGTTTATTTTGCCAATTATGGTTTATCTCA-AAATAGTTCA
SEQ1805	CGTATGGTC-CTAGACCTGCTCAACAGATTAGAGAGTTTACCTGGCAGATGGTTAACTT
SEQ1806	CGTATGGTC-CTAGACCTGCTCAACAGATTAGAGAGTTTACCTGGCAGATGGTTAATTT
SEQ1807	GATGACCACCGTGTTTATTTTGCCAATTATGGTTTATCTCA-AAATAGTTCA
SEQ1808	CGTATGGTC-CTAGACCTGCTCAACAGATTAGAGAGTTTACCTGGCAGATGGTTAATTT
SEQ1809	CGTATGGTC-CTAGACCTGCTCAACAGATTAGAGAGTTTACCTGGCAGATGGTTAACTT
SEQ1810	GATGACCACCGTGTTTATTTTGCCAATTATGGTTTATCTCA-AAATAGTTCA
SEQ1811	GATGACCACCGTGTTTATTTTGCCAATTATGGTTTATCTCA-AAATAGTTCA
SEQ1812	GATGACCACCGTGTTTATTTTGCCAATTATGGTTTATCTCA-AAATAGTTCA
SEQ1813	CGTATGGTC-CTAGACCTGCTCAACAGATTAGAGAGTTTACCTGGCAGATGGTTAACTT
SEQ1814	GATGACCACCGTGTTTATTTTGCCAATTATGGTTTATCTCA-AAATAGTTCA

Table 18: Comparative Sequences relating to SAG1600 (glutamate racemase)

SEQ1801	TTATTGACTAAAAATGTTAAGATGATTGTTATAGCTTGTAATACAGCAACTGCAGTTGC
SEQ1802	TTATTGACTAAAAATGTTAAGATGATTGTTATAGCTTGTAATACAGCAACTGCAGTTGC
SEQ1803	TTATTGACTAAAAATGTTAAGATGATTGTTATAGCTTGTAATACAGCAACTGCAGTTGC
SEQ1804	ATAAAACAGAAATATCACGAACGGT-TTCTGCGCCACTATCAATTAATTTAACCTCA
SEQ1805	TTATTGACTAAAAATGTTAAGATGATTGTTATAGCTTGTAATACAGCAACTGCAGTTGC
SEO1806	TTATTGACTAAAAATGTTAAGATGATTGTTATAGCTTGTAATACAGCAACTGCAGTTGC
SEQ1807	
	ATAAAACAGAAATATCACGAACGGT-TTCTGCGCCACTATCAATTAATTTAACCTCA
SEQ1808	TTATTGACTAAAAATGTTAAGATGATTGTTATAGCTTGTAATACAGCAACTGCAGTT
SEQ1809	TTATTGACTAAAAATGTTAAGATGATTGTTATAGCTTGTAATACAGCAACTGCAGTTGC
SEQ1810	ATAAAACAGAAATATCACGAACGGT-TTCTGCGCCACTATCAATTAATTTAACCTCA
SEQ1811	ATAAAACAGAAATATCACGAACGGT-TTCTGCGCCACTATCAATTAATTTAACCTCA
SEQ1812	ATAAAACAGAAATATCACGAACGGT-TTCTGCGCCACTATCAATTAATTTAACCTCA
SEQ1813	TTATTGACTAAAAATGTTAAGATGATTGTTATAGCTTGTAATACAGCAACTGC
SEQ1814	ATAAAACAGAAATATCACGAACGGT-TTCTGCGCCACTATCAATTAATTTAACCTCA
22223	initiation de la constitución de
GEO1 801	TGGCAAGAAATTAAAGAAAAACTAGACGTGCCTGTTTTAGGCGTTATTTTACCAGGAGC
SEQ1801	
SEQ1802	TGGCAAGAAATTAAAGAAAAACTAGACATCCCTGTTTTAGGCGTTATTTTACCAGGAGC
SEQ1803	TGGCAAGAAATTAAAGAAAAACTAGACATACCTGTTTTAGGCGTTATTTTACCAGGAGC
SEQ1804	CCCCCATAACATTTTGAATGATGGGACGTAATAGGGGATAATGC-GTGCAACCTAAAAT
SEQ1805	TGGCAAGAAATTAAAGAAAAACTAGACATCCCTGTTTTAGGCGTTATTTTACCAGGAGC
SEQ1806	TGGCAAGAAATTAAAGAAAAACTAGACATAC
SEQ1807	CCCCATAACATTTTGAATAATGGGACGTAATAGGGGATAATGC-GTGCAACCTAAAAT
SEQ1808	
SEQ1809	TGGCAAGAAATTAAAGAAAAACTAGACATCCCTGTTTTAGGCGTTATTTTACCAGGAGC
SEQ1810	
_	CCCCCATAACATTTTGAATGATGGGACGTAATATGGGATAATGC-GTGCAACCTAAAAT
SEQ1811	CCCCCATAACATTTTGAATGATGGGACGTAATAGGGGATAATGC-GTGCAACCTAAAAT
SEQ1812	CCCCCATAACATTTTGAATGATGGGACGTAATAGGGGATAATGC-GTGCAACCTAAAAT
SEQ1813	
SEQ1814	CCCCATAACATTTTGAATGATGGGACGTAATAAGGGATAATGC-GTGCAACCTAAAAT
SEQ1801	AGCGCAGCTATCAAATCAACTAATTCAGGGAAAGTTGGTATTATAGGTACTCCCATGAC
SEQ1802	AGCGCAGCTATCAAATCAACTAATTTAGGGAAAGTTGGTATTATAGGTACTCCCATGAC
SEQ1803	AGCGCAGCTATCAAATCAACTAATTCAGGGAAAGTTGGTATTATAGGTACTCCCATGAC
SEQ1804	AAAGTATCTAATTTACCAACTAATGGGGACAACGTTTCATAAACCACCTTTTTGGCTAA
SEQ1805	AGCGCAGCTATCAAATCAACTAATTTAGGGAAAGTTGGTATTATAGGTACTCCCATGAC
SEQ1806	AGGGGGG TATCAAATCAACTAATTTAGGGAAAGTTGGTATTATAGGTACTCCCATGAC
SEQ1807	AAAGTATCTAATTTACCAACTAATGGGGACAATGTTTCATAAACCACCTTTTTGGCTAA
SEQ1808	
SEQ1809	AGCGCAGCTATCAAATCAACTAATTTAGGGAAAGTTGGTATTATAGGTACTCCCATGAC
SEQ1810	AAAGTA
SEQ1811	AAAGTATCTAATTTACCAACTAATGGGGACAACGTTTCATAAACCACCTTTTTGGCTAA
SEQ1812	AAAGTATCTAATTTACCAACTAATGGGGACAACGTTTCATAAACCACCTTTTTGGCTAA
SEQ1813	
SE01814	AAAGTATCTAATTTACCAACTAATGGGGACAACGTTTCATAAACCACCTTTTTTGGCTAA
SEQIOIT	AMAGIATETAMTTIACCAMETAMTGGGGACAACGTTTCATAMACCACCTTTTTGGCTAA
CEO1901	
SEQ1801	GTTAAATCAGATGCTTATCGTCAAAAAATTCAAGCTTTGTCTCCAAATACTGCTGTGGT
SEQ1802	GTTAAATCAGATGCTTATCGTCAAAAAATTCAAGCTTTGTCTCCAAATACTGCTGTGGT
SEQ1803	GTTAAATCAGATGCTTATCGTCAAAAAATTCAAGCTTTGTCTCCAAATACTGCTGGT
SEQ1804	CTAGAAGACATCTGATTTGATTCCACAATTGGAACAAATTTCGGACAAGCAAG
SEQ1805	GTTAAATCAGATGCTTATCGTCAAAAAATTCAAGCTTTGTCTCCAAATACTGCTGTGGT
SEQ1806	
SEQ1807	CTAGAAGACATCTGATTTGATTCCACAATTGGAACAAATTTCGGACAAGCAAG
SEQ1808	
SEQ1809	GTTAAATCAGATGCTTATCGTCAAAAAATTCAAGC
SEQ1810	
SEQ1811	CTAGAAGACATCTGATTTGATTCCACAATTGGAACAA
SEQ1812	CTAGAAGA
SEQ1813	
SEQ1814	CTAGAAGACATCTGATTTGATTCCACAATTGGAACAAATTTCGGACAAGCAAG

# Table 18: Comparative Sequences relating to SAG1600 (glutamate racemase)

SEQ1801	TCCCTTGCTTGTCCGAAATTTGTTCCAATTGTGGAATCAAATCAGATGTCTTCTAGTTT
SEQ1802	TCCCTTGCTTGTCCGAAATTTGTTCCAATTGTGGAATCAAATCAGATGTCTTCTAGTTT
SEQ1803	TCCCTTGCTTGTCCGAAATTTGTTCCAATTGTGGAATCAAATCAGATGTCTTCTAGTTT
	ACACCA COM DEMOCRACIA CA DE COMPANA DE COMPANA CAGA I GILLI I CIAGITI
SEQ1804	ACAGCAGTATTTGGAGACAAAGCTTGAATTTTTTGACGATAAGCATCTGATTTAACAGT
SEQ1805	TCCCTTGCTTGTCCGAAAT
SEQ1806	
SEQ1807	ACAGCAGTATTTGGAGACAAAGCTTGAATTTTTTGACGATAAGCATCTGATTTAACAGT
SEQ1808	
SEQ1809	
SEQ1810	~
SEQ1811	
SEQ1812	
SEQ1813	
SEQ1814	ACAGCAGTATTTGGAGACAAAGCTTGAATTTTTTGACGATAAGCATCTGATTTAACAGT
_	
SEQ1801	GCCAAAAAGGTGGTTTATGAAACGTTGTCCCCATTAGTTGGTAAATTAGATACTTTAAT
SEQ1802	GCCAAAAAGGTGGTTTATGAAACGTTGTCCCCATTAGTTGGTAAATTAGATACTTTAAT
SEQ1803	GCCAAAAAGGTGGTTTATGAAACGCTGTCCCCATTAGTTGGTAAATTAGATACTTTAAT
SEQ1804	ATGGGAGTACCTATAATACCAACTTTCCCTAAATTAGTTGATTGA
SEQ1805	
SEQ1806	
SEQ1807	ATGGGAGTACCTATAA
SEQ1808	AIGGGAGIACCIAIAA
SEQ1809	
SEQ1810	
SEQ1811	
SEQ1812	
SEQ1813	
SEQ1814	ATGGGAGTACCTATAATACCAACTTTCCCTGAATABCMARATVSTNCSRATNGTSAGGT
SESTOTA	AIGGGAGIACCIAIAAIACCAACIIICCCIGAAIABCPARAIVSINCSRAINGISAGGI
SEQ1801	TTAGGTTGCACGCATTATCCCTTATTACGTCCCATCATTCAAAATGTTATGGGGGCTGA
SEQ1802	TTAGGTTGCACGCATTATCCCCTATTACGTCCCATCATTCAAAATGTTATGGGGGCTGA
SEQ1803	TTAGGTTGCACGCATTATCCCTTATTACGTCCCATCATTCAAAATGTTATGGGGGCTGA
SEQ1804	CCTGGTAAAATAACGCCTAAAACAGGGATGTCTAGTTTTTCTTTAATTTCTTGCCAGGC
SEQ1805	
SEQ1806	
SEQ1807	
SEQ1808	
SEQ1809	
SEQ1810	
SEQ1811	
SEQ1812	
SEQ1813	
SEQ1013 SEO1814	
SEQ1614	AMATRACMAS
anot 001	
SEQ1801	GTTAAATTAATTGATAGTGGCGCAGAAACCGTTCGTGATATTTCTGTTTTATTGAACTA
SEQ1802	GTTAAATTAATTGATAGTGGCGCAGAAACCGTTCGTGATATTTCTGTTTTATTGAACTA
SEQ1803	GTTAAATTAATTGATAGTGGCGCAGAAACCGTTCGTGATATTTCTGTTTTATTGAACTA
SEQ1804	ACTGCAGTTGCTGTATTACAAGCTATAACAATCATCTTAACATTTTTAGTCAATAAGAA
SEQ1805	
SEO1806	
	·
SEQ1807	
SEQ1808	
SEQ1809	
SEQ1810	
SEQ1811	
SEQ1812	
SEQ1813	
SEQ1814	

# Table 18: Comparativ Sequences relating to SAG1600 (glutamate racemase)

SEQ1801	TTTGAGATAAACCATAATTGGCAAAATAAACACGGTGGTCATCACTTTTACACAACCGC
SEQ1802	TTTGAGATAAACCATAATTGGCAAAATAAACACGGTGGTCATCACTTTTACACAACCGC
SEQ1803	TTTGAGATAAMCCATAATTGGSMAAATAAACACGGTGGTCATCACTTTTACACAACCGS
SEQ1804	TTAACCATCTGCCAGGTAAACTCTCTAATCTGTTGAGCAGGTCTAGGACCATACGGAGC
SEQ1805	
SEQ1806	
SEQ1807	
SEQ1808	
SEQ1809	
SEQ1810	
SEQ1811	
SEQ1812 .	
SEQ1813	
SEQ1814	
•	
SEQ1801	AGCCCAA
SEQ1802 ·	AGCCCAAAAGGTTTTAAAGAAA
SEQ1803	AGCCCAAAAGGTTTTTAAGGAAATTGCAGAACAATGGCTTAATCAAGAAATAAAT
SEQ1804	CTAGCCTGATCTCCAATGAAGATTACTTCCTCTTCTGGAAGTTGACGGAACATTTCCTT
SEQ1805	
SEQ1806	· •••••
SEQ1807	
SEQ1808	
SEQ1809	
SEQ1810	
SEQ1811	
SEQ1812	
SEQ1813	
SEQ1814	
_	•
SEQ1801	
SEQ1802	
SEQ1803	
SEQ1804	ACAACCGTTAAACCACCT
SEQ1805	~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~
SEQ1806	*
SEQ1807	
SEQ1808	
SEQ1809	
SEQ1810	
SEQ1811	
SEQ1812	
SEQ1813	~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~
SEQ1814	

### Table 19: C mparative Sequences relating to SAF1680 (shikimate 5-dehydrogenase)



## Table 19: Comparative Sequences relating to SAF1680 (shikimate 5-dehydr genase)

SEQ ID NO. 1907: SAG1680 FROM THE COH1 GBS TYPE Ia STRAIN
TGCACGCCACTCTCTATCCCCGTTAATGTGGAATACCTCTTTTAAGAAAAAAACATGAATTATGCCTATCTGACATTTGA
AGTAGAAGAGGGGTAAATTAACAGAAGCTGTTCGAGGTGTCAGGGCATTGAGTATTCGTGGTGTTAATGTTTCAATGCCAT
TTAAACAGAGTGTTATCCCTTTGCTAGATGATTTATCTCCTCAAGCTAAATTAGTGGGTGCTGTAAATACT

SEQ ID NO. 1908: SAG1680 FROM THE CJB110 GBS NONTYPEABLE STRAIN
ATTCGTTATTAATTGAAATGCTTCTGCTCCTTGATAAATCAGCATCCCTAGACCATTATAAAGCATGTTTCACTCCATTTT
GTCTAACAAATCGTAACAATGCTGTTTCTTTAGGCTTGTAAACCAAGTCGACAACTACTAAATTGGGTGTTAAAATTTCT
GGATCGTTAATTAAACTATAATTATCTAATGGCCTCATTCCTAAACTAGTAGCATCAATATAAAAATGACTAGTTCTAAT
AGCGTCTTTAAATGCTGTCTTATTTTCTAGATAATCAACGACTACCTTTATTTGAAACTGTTTTTTAATTTTATCTGATA
AGTCAATGACCTTATCGTAATTTGAGCTATTACGATTAAATAATCTÄATTTCCGCAACTCCCTCCATAACTGCTTGAACT
GCAACTGCTTTACCTGAACCACCAATACCAGCTATTGTAACTATTTT

SEQ ID NO. 1909: SAG1680 FROM THE CJB110 GBS NONTYPEABLE STRAIN
ACTCTCTATCCCCGTTAATGTGGAATACCTCTTTTCAAGAAAAAACATGAATTATGCCTATCTGACATTTGAAGTAGAA
GAGGGTAAATTAACAGAAGCTGTTCGAGGTGTCAGGGCATTGGGTATTCGTGGTGTTAATGTTTCAATGCCATTTAAACA
GAGTGTTATCCCTTTGCTAGATGATTTATCTCCTCAAGCTAAATTAGTGGGTGCTGTAAATACTATCGTTAATCAAGGTG
GAACCGGACGTTTAGTAGGCCATATGACAGATGGCATTGGTTGTTTTAAAGCTTTAGCAGCTCAAGGTTTCAGTGCTAAA
AATAAAATAGTTACAATAGCTGGTATTGGTG

SEQ ID NO. 1910: SAG1680 FROM THE 1169NT1 GBS TYPE V STRAIN
ATTCGTTATTAATTGAAATGCTTCTGCTCCTTGATAAATCAGCATCCCTAGACCATTATAAGCATGTTTCACTCCATTTT
GTCTAACAAATCGTAACAATGCTGTTTCTTTAGGCTTGTAAACCAAGTCGACAACTACTAAATTCGGTGTTAAAATTTCT
GGATCGTTAATTAAACTATAATTATCTAATGGCCTCATTCCTAAACTAGTAGCATCAATATAAAAATGACTAGTTCTAAT
AGCGTCTTTAAATGCTGTCTTATTTTCTAGATAATCAACGACTACCTTTATTTGAAACTGTTTTTTAATTTTATCTGATA
AGTCAATGACCTTATCGTAATTTGAGCTGTTACGAT

SEQ ID NO. 1911: SAG1680 FROM THE 1169NT1 GBS TYPE V STRAIN
ACTTCTCTATTCCCCGTTAATGTGGAATACCTCTTTTCAAGAAAAAACATGAATTATGCCTATCTGACATTTGAAGTAG
AAGAGGGTAAATTAACAGAAGCTGTTCGAGGTGTCAGGGCATTGGGTATTCGTGGTGTTAATGTTTCAATGCCATTTAAA
CAGAGTGTTATCCCTTTGCTAGATGATTTATCTCCTCAAGCTAAATTAGTGGGTGCTGTAAATACTATCGTTAATCAAGG
TGGAACC

SEQ ID NO. 1912: SAG1680 FROM THE 18RS21 GBS TYPE II STRAIN
TCGTTATTAATTGAAATGCTTCTGCTCCTTGATAAATCATCATCCCTAGACCATTATAAGCATGTTTCACTCCATTTTGT
CTAACAAATCGTAACAATGCTGTTTCTTTAGGCTTGTAAACCAAGTCGACAACTACTAAATTCGGTGTTAAAATTTCTGG
ATCGTTAATTAAACTATAATTATCTAATGGCCTCATTCCTAAACTAGTAGCATCAATATAAAAATGACTAGTTCTAATAG
CGTCTTTAAATGCTGTCTTATTTTCTAGATAATCAACGACTACCTTTATTTGAAACTGTTTTTTAATTTTATCTGATAAG
TCAATGACCTTATCGTAATTTGAGCTGTTACGATTAAATAATCTAATTTCCGCAAC

## Table 19: C mparative Sequences relating t SAF1680 (shikimate 5-dehydrogenase)

SEQ1902 TGGTCTAATGCCAATCCTGCACGCCACTCTCTAT—CCCCGTTAAATGCGAATCCCCT SEQ1904 ————————————————————————————————————
SEQ1903       TGGTCTAATTGCCAATCCTGCACGCCACTCTCTAT-CCCCGTTAATGTGAATACCTCT         SEQ1904       ————————————————————————————————————
SEQ1904 SEQ1905 SEQ1906 SEQ1907 SEQ1908 SEQ1908 SEQ1909 SEQ1909 SEQ1909 SEQ1910 SEQ1910 SEQ1910 SEQ1910 SEQ1910 SEQ1910 SEQ1910 SEQ1911 SEQ1911 SEQ1912 SEQ1912 SEQ1913 SEQ1914 SEQ1914 SEQ1914 SEQ1901 GACCATTATAAG—CATGTTTCACTCCATTTTGTCTAACAAATCGTAACAATGCTGTTTCSEQ1914 SEQ1910 SEQ1911 SEQ1911 SEQ1912 SEQ1913 SEQ1914 SEQ1901 SEQ1901 SEQ1901 SEQ1901 SEQ1901 SEQ1901 SEQ1901 SEQ1902 GACCATTATAAG—CATGTTTCACTCCATTTTGTCTAACAAATCGTAACAATGCTGTTTCCTCTCTCT
SEQ1906TGCACGCCACTCTCTAT-CCCCGTTAATGTGAATACCTCT SEQ1908ATTCGTTATTAATTGAAATGCTTCTGCTCCTTGATAAATCAGCATCCCT SEQ1909ACTCTCTAT-CCCCGTTAATGTGGAATACCTCT SEQ1910ACTCTCTAT-CCCCGTTAATGTGGAATACCTCT SEQ1911
SEQ1907TGCACGCCACTCTCTAT-CCCCGTTAATGTGGAATACCTCT SEQ1908ATTCGTTATTAATTGAAATGCTTCTGCTCCTTGATAAATCAGCATCCCT SEQ1909
SEQ1907TGCACGCCACTCTCTAT-CCCCGTTAATGTGGAATACCTCT SEQ1908ATTCGTTATTAATTGAAATGCTTCTGCTCCTTGATAAATCAGCATCCCT SEQ1909
SEQ1909
SEQ1910 SEQ1911 SEQ1911 SEQ1912 SEQ1913 SEQ1914 SEQ1914 SEQ1914 SEQ1901 GACCATTATAAG—CATGTTTCACTCCATTTTGTCTAACAAATCGTAACAATGCTGTTTC SEQ1902 SEQ1903 SEQ1903 TTCAAGAAAAAAACATGAATTATGCCTATCTGACAATTGAACAATGCTGTTTC SEQ1904 SEQ1905 SEQ1906 SEQ1907 SEQ1907 SEQ1907 SEQ1908 GACCATTATAAG—CATGTTTCACTCCATTTTGTCTAACAAATCGTAACAATGCTGTTTC SEQ1909 SEQ1901 SEQ1901 SEQ1901 SEQ1901 SEQ1902 GACCATTATAAG—CATGTTTCACTCCATTTTGTCTAACAAATCGTAACAATGCTGTTTC SEQ1903 SEQ1904 SEQ1905 SEQ1906 SEQ1906 SEQ1907 SEQ1907 SEQ1907 SEQ1907 SEQ1907 SEQ1908 SACCATTATAAG—CATGTTTCACTCCATTTTGTCTAACAAATCGTAACAATGCTGTTTC SEQ1907 SEQ1908 SACCATTATAAG—CATGTTTCACTCCATTTTGTCTAACAAATCGTAACAATGCTGTTTC SEQ1909 SEQ1910 GACCATTATAAG—CATGTTTCACTCCATTTTGTCTAACAAATCGTAACAATGCTGTTTC SEQ1910 SEQ1911 SEQ1911 GACCATTATAAG—CATGTTTCACTCCATTTTGTCTAACAAATCGTAACAATGCTGTTTC SEQ1911 TTCAAGAAAAAACATGAATTATGCCTATCTGACATTTGAAGAAGAGGGTAAATTA SEQ1911 SEQ1912 GACCATTATAAG—CATGTTTCACTCCATTTTGTCTAACAAATCGTAACAATGCTGTTTC SEQ1912 GACCATTATAAG—CATGTTTCACTCCATTTTGTCTAACAAATCGTAACAATGCTGTTTC SEQ1913
SEQ1910ATTCGTTATTAATTGAAATGCTTCTGCTCCTTGATAAATCAGCATCCCT SEQ1911
SEQ1912 SEQ1913 SEQ1914 SEQ1901 GACCATTATAAG—CATGTTTCACTCCATTTTGTCTAACAAATCGTAACAATGCTGTTTC SEQ1902 SEQ1903 TTCAAGAAAAAAACATGATTTCACTCCATTTTGTCTAACAAATCGTAACAATGCTGTTTC SEQ1904 GACCATTATAAG—CATGTTTCACTCCATTTTGTCTAACAAATCGTAACAATGCTGTTTC SEQ1905 SEQ1906 GACCATTATAAG—CATGTTTCACTCCATTTTGTCTAACAAATCGTAACAATGCTGTTTC SEQ1907 TT-AAGAAAAAAACATGAATTATGCCTATCTGACATTTGAAGTAGAAGAGGGTAAATTA SEQ1908 SEQ1909 GACCATTATAAG—CATGTTTCACTCCATTTTGTCTAACAAATCGTAACAATGCTGTTTC SEQ1909 TT-AAGAAAAAAACATGAATTATGCCTATCTGACATTTGAAGTAGAAGAGGGTAAATTA SEQ1908 GACCATTATAAG—CATGTTTCACTCCATTTTGTCTAACAAATCGTAACAATGCTGTTTC SEQ1909 TTCAAGAAAAAAACATGAATTATGCCTATCTGACATTTGAAGTAGAAGAGGGTAAATTA SEQ1910 GACCATTATAAG—CATGTTTCACTCCATTTTGTCTAACAAATCGTAACAATGCTGTTTC SEQ1910 GACCATTATAAG—CATGTTTCACTCCATTTTGTCTAACAAATCGTAACAATGCTGTTTC SEQ1911 TTCAAGAAAAAAACATGAATTATGCCTATCTGACATTTGAAGTAGAAGAGGGTAAATTA SEQ1912 GACCATTATAAG—CATGTTTCACTCCATTTTGTCTAACAAATCGTAACAATGCTGTTTC SEQ1912 GACCATTATAAG—CATGTTTCACTCCATTTTGTCTAACAAATCGTAACAATGCTGTTTC SEQ1913
SEQ1912 SEQ1913 SEQ1914 SEQ1901 GACCATTATAAG-CATGTTTCACTCCATTTTGTCTAACAAATCGTAACAATGCTGTTTC SEQ1902 GACCATTATAAG-CATGTTTCACTCCATTTTGTCTAACAAATCGTAACAATGCTGTTTC SEQ1903 TTCAAGAAAAAACATGAATTATGCCTATCTGACATTTGAAGAAGAGGGTAAATTA SEQ1904 GAC-ATTATAAG-CATGTTTCACTCCATTTTGTCTAACAAATCGTAACAATGCTGTTTC SEQ1905 SEQ1906 GACCATTATAAT-CATGTTTCACTCCATTTTGTCTAACAAATCGTAACAATGCTGTTTC SEQ1907 TT-AAGAAAAAAACATGAATTATGCCTATCTGACATTTGAAGTAGAAGAGGGTAAATTA SEQ1908 GACCATTATAAG-CATGTTTCACTCCATTTTGTCTAACAAATCGTAACAATGCTGTTTC SEQ1909 TTCAAGAAAAAAACATGAATTATGCCTATCTGACATTTGAAGTAGAAGAGGGTAAATTA SEQ1910 GACCATTATAAG-CATGTTTCACTCCATTTTGTCTAACAAATCGTAACAATGCTGTTTC SEQ1910 GACCATTATAAG-CATGTTTCACTCCATTTTGTCTAACAAATCGTAACAATGCTGTTTC SEQ1911 TTCAAGAAAAAAACATGAATTATGCCTATCTGACATTTGAAGTAGAAGAGGGTAAATTA SEQ1912 GACCATTATAAG-CATGTTTCACTCCATTTTGTCTAACAAATCGTAACAAATCGTGTTTC SEQ1912 GACCATTATAAG-CATGTTTCACTCCATTTTGTCTAACAAATCGTAACAAATCGTGTTTC SEQ1913
SEQ1914  SEQ1914  GACCATTATAAG-CATGTTTCACTCCATTTTGTCTAACAAATCGTAACAATGCTGTTTC SEQ1902  GACCATTATAAG-CATGTTTCACTCCATTTTGTCTAACAAATCGTAACAATGCTGTTTC SEQ1903  TTCAAGAAAAAAACATGAATTATGCCTATCTGACATTTGAAGTAGAAGAGGGTAAATTA SEQ1904  GAC-ATTATAAG-CATGTTTCACTCCATTTTGTCTAACAAATCGTAACAATGCTGTTTC SEQ1905  SEQ1906  GACCATTATAAT-CATGTTTCACTCCATTTTGTCTAACAAATCGTAACAATGCTGTTTC SEQ1907  TT-AAGAAAAAAACATGAATTATGCCTATCTGACATTTGAAGTAGAAGAGGGTAAATTA SEQ1908  GACCATTATAAG-CATGTTTCACTCCATTTTGTCTAACAAATCGTAACAATGCTGTTTC SEQ1909  TTCAAGAAAAAAACATGAATTATGCCTATCTGACATTTGAAGTAGAAGAGGGTAAATTA SEQ1910  GACCATTATAAG-CATGTTTCACTCCATTTTGTCTAACAAATCGTAACAATGCTGTTTC SEQ1911  TTCAAGAAAAAAACATGAATTATGCCTATCTGACATTTGAAGTAGAAGAGGGTAAATTA SEQ1912  GACCATTATAAG-CATGTTTCACTCCATTTTGTCTAACAAATCGTAACAAATCGTTTC SEQ1913
SEQ1914  SEQ1901  GACCATTATAAG-CATGTTTCACTCCATTTTGTCTAACAAATCGTAACAATGCTGTTTC SEQ1902  GACCATTATAAG-CATGTTTCACTCCATTTTGTCTAACAAATCGTAACAATGCTGTTTC SEQ1903  TTCAAGAAAAAACATGAATTATGCCTATCTGACATTTGAAGTAGAAGAGGGTAAATTA SEQ1904  GAC-ATTATAAG-CATGTTTCACTCCATTTTGTCTAACAAATCGTAACAATGCTGTTTC SEQ1905  SEQ1906  GACCATTATAAT-CATGTTTCACTCCATTTTGTCTAACAAATCGTAACAATGCTGTTTC SEQ1907  TT-AAGAAAAAAACATGAATTATGCCTATCTGACATTTGAAGTAGAAGAGGGTAAATTA SEQ1908  GACCATTATAAG-CATGTTTCACTCCATTTTGTCTAACAAATCGTAACAATGCTGTTTC SEQ1909  TTCAAGAAAAAAACATGAATTATGCCTATCTGACATTTGAAGTAGAAGAGGGTAAATTA SEQ1910  GACCATTATAAG-CATGTTTCACTCCATTTTGTCTAACAAATCGTAACAATGCTGTTTC SEQ1911  TTCAAGAAAAAAACATGAATTATGCCTATCTGACATTTGAAGTAGAAGAGGGTAAATTA SEQ1912  GACCATTATAAG-CATGTTTCACTCCATTTTGTCTAACAAATCGTAACAATGCTGTTTC SEQ1913
SEQ1902 GACCATTATAAG-CATGTTTCACTCCATTTTGTCTAACAAATCGTAACAATGCTGTTTC SEQ1903 TTCAAGAAAAAACATGAATTATGCCTATCTGACATTTGAAGTAGAAGAGGGTAAATTA SEQ1904 GAC-ATTATAAG-CATGTTTCACTCCATTTTGTCTAACAAATCGTAACAATGCTGTTTC SEQ1905
TTCAAGAAAAAACATGAATTATGCCTATCTGACATTTGAAGTAGAAGAGGGGTAAATTA SEQ1904 GAC-ATTATAAG-CATGTTTCACTCCATTTTGTCTAACAAATCGTAACAATGCTGTTTC SEQ1905 SEQ1906 GACCATTATAAT-CATGTTTCACTCCATTTTGTCTAACAAATCGTAACAATGCTGTTTC SEQ1907 TT-AAGAAAAAACATGAATTATGCCTATCTGACATTTGAAGTAGAAGAGGGTAAATTA SEQ1908 GACCATTATAAG-CATGTTTCACTCCATTTTGTCTAACAAATCGTAACAATGCTGTTTC SEQ1909 TTCAAGAAAAAAACATGAATTATGCCTATCTGACATTTGAAGTAGAAGAGGGTAAATTA SEQ1910 GACCATTATAAG-CATGTTTCACTCCATTTTGTCTAACAAATCGTAACAATGCTGTTTC SEQ1911 TTCAAGAAAAAAACATGAATTATGCCTATCTGACATTTGAAGTAGAAGAGGGTAAATTA SEQ1912 GACCATTATAAG-CATGTTTCACTCCATTTTGTCTAACAAATCGTAACAATGCTGTTTC SEQ1913
SEQ1904 GAC-ATTATAAG-CATGTTTCACTCCATTTTGTCTAACAAATCGTAACAATGCTGTTTC SEQ1905 SEQ1906 GACCATTATAAT-CATGTTTCACTCCATTTTGTCTAACAAATCGTAACAATGCTGTTTC SEQ1907 TT-AAGAAAAAACATGAATTATGCCTATCTGACATTTGAAGTAGAAGAGGGTAAATTA SEQ1908 GACCATTATAAG-CATGTTTCACTCCATTTTGTCTAACAAATCGTAACAATGCTGTTTC SEQ1909 TTCAAGAAAAAAACATGAATTATGCCTATCTGACATTTGAAGTAGAAGAGGGTAAATTA SEQ1910 GACCATTATAAG-CATGTTTCACTCCATTTTGTCTAACAAATCGTAACAATGCTGTTTC SEQ1911 TTCAAGAAAAAAACATGAATTATGCCTATCTGACATTTGAAGTAGAAGAGGGTAAATTA SEQ1912 GACCATTATAAG-CATGTTTCACTCCATTTTGTCTAACAAATCGTAACAATGCTGTTTC SEQ1913
SEQ1905 SEQ1906 GACCATTATAAT-CATGTTTCACTCCATTTTGTCTAACAAATCGTAACAATGCTGTTTC SEQ1907 TT-AAGAAAAAACATGAATTATGCCTATCTGACATTTGAAGTAGAAGAGGGTAAATTA SEQ1908 GACCATTATAAG-CATGTTTCACTCCATTTTGTCTAACAAATCGTAACAATGCTGTTTC SEQ1909 TTCAAGAAAAAAACATGAATTATGCCTATCTGACATTTGAAGTAGAAGAGGGTAAATTA SEQ1910 GACCATTATAAG-CATGTTTCACTCCATTTTGTCTAACAAATCGTAACAATGCTGTTTC SEQ1911 TTCAAGAAAAAAACATGAATTATGCCTATCTGACATTTGAAGTAGAAGAGGGTAAATTA SEQ1912 GACCATTATAAG-CATGTTTCACTCCATTTTGTCTAACAAATCGTAACAATGCTGTTTC SEQ1913
SEQ1906 GACCATTATAAT-CATGTTTCACTCCATTTTGTCTAACAAATCGTAACAATGCTGTTTC SEQ1907 TT-AAGAAAAAACATGAATTATGCCTATCTGACATTTGAAGTAGAAGAGGGTAAATTA SEQ1908 GACCATTATAAG-CATGTTTCACTCCATTTTGTCTAACAAATCGTAACAATGCTGTTTC SEQ1909 TTCAAGAAAAAAACATGAATTATGCCTATCTGACATTTGAAGTAGAAGAGGGTAAATTA SEQ1910 GACCATTATAAG-CATGTTTCACTCCATTTTGTCTAACAAATCGTAACAAATCGTTTC SEQ1911 TTCAAGAAAAAAACATGAATTATGCCTATCTGACATTTGAAGTAGAAGAGGGTAAATTA SEQ1912 GACCATTATAAG-CATGTTTCACTCCATTTTGTCTAACAAATCGTAACAATGCTGTTTC SEQ1913
SEQ1907 TT-AAGAAAAAACATGAATTATGCCTATCTGACATTTGAAGTAGAAGAGGGTAAATTA SEQ1908 GACCATTATAAG-CATGTTTCACTCCATTTTGTCTAACAAATCGTAACAAATCGTTTC SEQ1909 TTCAAGAAAAAAACATGAATTATGCCTATCTGACATTTGAAGTAGAAGAGGGTAAATTA SEQ1910 GACCATTATAAG-CATGTTTCACTCCATTTTGTCTAACAAATCGTAACAATGCTGTTTC SEQ1911 TTCAAGAAAAAAACATGAATTATGCCTATCTGACATTTGAAGTAGAAGAGGGTAAATTA SEQ1912 GACCATTATAAG-CATGTTTCACTCCATTTTGTCTAACAAATCGTAACAATGCTGTTTC SEQ1913
SEQ1908 GACCATTATAAG-CATGTTTCACTCCATTTTGTCTAACAAATCGTAACAATGCTGTTTC SEQ1909 TTCAAGAAAAAACATGAATTATGCCTATCTGACATTTGAAGTAGAAGAGGGTAAATTA SEQ1910 GACCATTATAAG-CATGTTTCACTCCATTTTGTCTAACAAATCGTAACAATGCTGTTTC SEQ1911 TTCAAGAAAAAACATGAATTATGCCTATCTGACATTTGAAGTAGAAGAGGGGTAAATTA SEQ1912 GACCATTATAAG-CATGTTTCACTCCATTTTGTCTAACAAATCGTAACAATGCTGTTTC SEQ1913ATGCCTATCTGACATTTGAAGTAGAAGAGGGTAAATTA
SEQ1909 TTCAAGAAAAAACATGAATTATGCCTATCTGACATTTGAAGTAGAAGAGGGGTAAATTA SEQ1910 GACCATTATAAG-CATGTTTCACTCCATTTTGTCTAACAAATCGTAACAATGCTGTTTC SEQ1911 TTCAAGAAAAAAACATGAATTATGCCTATCTGACATTTGAAGTAGAAGAGGGGTAAATTA SEQ1912 GACCATTATAAG-CATGTTTCACTCCATTTTGTCTAACAAATCGTAACAATGCTGTTTC SEQ1913ATGCCTATCTGACATTTGAAGTAGAAGAGGGTAAATTA
SEQ1910 GACCATTATAAG-CATGTTTCACTCCATTTTGTCTAACAAATCGTAACAATGCTGTTTC SEQ1911 TTCAAGAAAAAACATGAATTATGCCTATCTGACATTTGAAGTAGAAGAGGGTAAATTA SEQ1912 GACCATTATAAG-CATGTTTCACTCCATTTTGTCTAACAAATCGTAACAATGCTGTTTC SEQ1913ATGCCTATCTGACATTTGAAGTAGAAGAGGGTAAATTA
SEQ1911 TTCAAGAAAAAACATGAATTATGCCTATCTGACATTTGAAGTAGAAGAGGGTAAATTA SEQ1912 GACCATTATAAG-CATGTTTCACTCCATTTTGTCTAACAAATCGTAACAATGCTGTTTC SEQ1913ATGCCTATCTGACATTTGAAGTAGAAGAGGGTAAATTA
SEQ1912 GACCATTATAAG-CATGTTTCACTCCATTTTGTCTAACAAATCGTAACAATGCTGTTTC SEQ1913ATGCCTATCTGACATTTGAAGAAGAGAGGGTAAATTA
SEQ1913ATGCCTATCTGACATTTGAAGTAGAAGAGGGTAAATTA
•
SEO1 91 4 GACCAጥጥሕጥሕ በሚያቸርች የሚያቸው የ
GIOGRIPHENGIONIGIIIONGICONIIIIGICIMOMMIGGIMOMIGGIGIII
SEQ1901 TTAGGCTTGTAAACCAAGTCGACAACTACTAAATTCGGTGTTAAAATTTCTGGATCG
SEQ1902 TTAGGCTTGTAAACCAAGTCGACAACTACTAAATTCGGTGTTAAAATTTCTGGATCG
SEQ1903 CAGAAGCTGTTCGAGGTGTCAGGGCATTGAGTATTCGTGGTGTTAATGTTTCAATGCCA
SEQ1904 TTAGGCTTGTAAACCAAGTCGACAACTACTAAATTCGGTGTTAAAATTTCTGGATCC
SEQ1905GTTCGAGGTGTCAGGGCATTGGGTATTCGTGGTGTTAATGTTTCAATGCCA
SEQ1906 TTAGGCTTGTAAACCAAGTCGACAACTACTAAATTCGGTGTTAAAATTTCTGGATCC
SEQ1907 CAGAAGCTGTTCGAGGTGTCAGGGCATTGAGTATTCGTGGTGTTAATGTTTCAATGCCA
SEQ1908 TTAGGCTTGTAAACCAAGTCGACAACTACTAAATTGGGTGTTAAAATTTCTGGATCC
SEQ1909 CAGAAGCTGTTCGAGGTGTCAGGGCATTGGGTATTCGTGGTGTTAATGTTTCAATGCCA
SEQ1910 TTAGGCTTGTAAACCAAGTCGACAACTACTAAATTCGGTGTTAAAATTTCTGGATC
SEQ1911 CAGAAGCTGTTCGAGGTGTCAGGGCATTGGGTATTCGTGGTGTTAATGTTTCAATGCC
SEQ1912 TTAGGCTTGTAAACCAAGTCGACAACTACTAAATTCGGTGTTAAAATTTCTGGATC
SEQ1913 CAGAAGCTGTTCGAGGTGTCAGGGCATTGGGTATTCGTGGTGTTAATGTTTCAATGCC
SEQ1914 TTAGGCTTGTAAACCAAGTCGACAACTACTAAATTGGGTGTTAAAATTTCTGGATC

## Table 19: Comparative Sequences relating t SAF1680 (shikimate 5-dehydrogenase)

SEQ1901	TT-AATTAAACTATAATTATCTAATGGCCTCATTCCT-AAACTAGTAGCATCAAT
SEQ1902	TT-AATTAAACTATAATTATCTAATGGCCTCATTCCT-AAACTAGTAGCATCAAT
SEQ1903	TTTAAACAGAGTGTTATCCCTTTGCTAGATGATTTATCTCCTCAAGCTAAATTAGTGGGT
SEQ1904	TT-AATTAAACTATAATTATCTAATGGCCTCATTCCT-AAACTAGTAGCATCAAT
SEQ1905	TTTAAACAGAGTGTTATCCCTTTGCTARATGATTTATCTCCTCAAGCTAAATTAGTGGGT
SEQ1906	TT-AATTAAACTATAATTATCTAATGGCCTCATTCCT-AAACTAGTAGCATCAAT
SEQ1907	TTTAAACAGAGTGTTATCCCTTTGCTAGATGATTTATCTCCTCAAGCTAAATTAGTGGGT
SEQ1908	TT-AATTAAACTATAATTATCTAATGGCCTCATTCCT-AAACTAGTAGCATCAAT
SEQ1909	TTTAAACAGAGTGTTATCCCTTTGCTAGATGATTTATCTCCTCAAGCTAAATTAGTGGGT
SEQ1910	TT-AATTAAACTATAATTATCTAATGGCCTCATTCCT-AAACTAGTAGCATCAAT
SEQ1911	TTTAAACAGAGTGTTATCCCTTTGCTAGATGATTTATCTCCTCAAGCTAAATTAGTGGGT
SEQ1912	TT-AATTAAACTATAATTATCTAATGGCCTCATTCCT-AAACTAGTAGCATCAAT
SEQ1913	TTTAAACAGAGTGTTATCCCTTTGCTAGATGATTTATCTCCTCAAGCTAAATTAGTGGGT
SEQ1914	TT-AATTAAACTATAATTATCTAATGGCCTCATTCCT-AAACTAGTAGCATCAAT
- <b>-</b>	
SEQ1901	TAAAAATGACTAGTTCTAATAGCGTCTTTAAATGCTGTCTTATTTTCTAGATAATCAAC
SEQ1902	TAAAAATGACTAGTTCTAATAGCGTCTTTAAATGCTGTCTTATTTTCTAGATAATCAAC
SEQ1903	CTGTAAATACTATCGTTAATCAAGGTGGAACCGGACGTTTAGTAGGCCATATGACAGAT
SEQ1904	TAAAAATGACTAGTTCTAATAGCGTCTTTAAATGCTGTCTTATTTTCTAGATAATCAAC
SEQ1905	CTGTAAATACTATCGTTAATCAAGGTGGAACCGSACGTTTAGTAGGCCATATGACAGAT
SEQ1906	TAAAAATGACTAGTTCTAATAGCGTCTTTAAATGCTGTCTTATTTTCTAGATAATCAAC
SEQ1907	CTGTAAATACT
SEQ1907	TAAAAATGACTAGTTCTAATAGCGTCTTTAAATGCTGTCTTATTTTCTAGATAATCAAC
SEQ1908	CTGTAAATACTATCGTTAATCAAGGTGGAACCGGACGTTTAGTAGGCCATATGACAGAT
SEQ1909 SEQ1910	TAAAAATGACTAGTTCTAATAGCGTCTTTAAATGCTGTCTTATTTTCTAGATAATCAAC
_	CTGTAAATGACTAGTTCTAATCAAGGTGGAACC
SEQ1911 SEQ1912	TAAAAATGACTAGTTCTAATAGCGTCTTTAAATGCTGTCTTATTTTCTAGATAATCAAC
SEQ1912 SEQ1913	CTGTAAATGCTAGTTCTAATGCGTCTTTAATGCTGTCTTATTTTCTAGTAGTCAAC
SEQ1913 SEQ1914	TAAAAATGACTACGTTAATAGCGTCTTTAAATGCTGTCTTATTTTCTAGATAATCAAC
2FÖT3T4	TAAAAATGACTAGTTCTAATAGCGTCTTTAATTCTAGATAATCAAC
SEQ1901	ACTACCTTTATTTGAAACTGTTTTTTAATTTTATCTGATAAGTCAATGACCTTATCGTA
SEQ1901 SEQ1902	ACTACCTTTATTTGAAACTGTTTTTTAATTTTATCTGATAAGTCAATGACCTTATCGTA
SEQ1903	GCATTGGTTGTTTTAAAGCTTTTAGCAGCTCAAGGTTTCAGTGCTAAAAATAAAT
SEQ1903 SEQ1904	ACTACCTTTATTTGAAACTGTTTTTTAATTTTATCTGATAAGTCAATGACCTTATCGTA
SEQ1905	GCATTGGTTGTTTTAAAGCTTTTAGCAGCTCAAGGTTTCAGTGCTAAAAATAAAT
SEQ1905	ACTACCTTTATTTGAAACTGTTTTTTAATTTTATCTGATAAGTCAATGACCTTATCGTA
SEQ1907	ACIACCITATIONACIGITITIANITIATIONACIANICACOTATORIA
<del></del>	ACTACCTTTATTTGAAACTGTTTTTTAATTTTATCTGATAAGTCAATGACCTTATCGTA
SEQ1908 SEQ1909	GCATTGGTTTTTTAAAGCTTTTAGCAGCTCAAGGTTTCAGTGCTAAAAATAAAATAGTT
_	ACTACCTTTATTTGAAACTGTTTTTTAATTTTATCTGATAAGTCAATGACCTTATCGTA
SEQ1910	ACTACCTTTATTTGAAACTGTTTTTTAATTTTATCTGATAAGTCAATGACCTTATCGT
SEQ1911	
SEQ1912	ACTACCTTTATTTGAAACTGTTTTTTAATTTTATCTGATAAGTCAATGACCTTATCGTA
SEQ1913	GCATTGGTTGTTTTAAAGCTTTAGCAGCTCAAGGTTTCAGTGCTAAAAATAAAATAAAT
SEQ1914	ACTACCTTTATTTGAAACTGTTTTTTAATTTTATCTGATAAGTCAATGACCTTATCGTA
GT-01-001	™™™™™™™™™™™™™™™™™™™™™™™™™™™™™™™™™™™™™
SEQ1901	TTTGAGCTGTTACGATTAAATAATCTAATTTCCGCAACTCCCTCC
SEQ1902	TTTGAGCTGTTACGATTAAATAATCTAATTTCCGCAACTCCCTCC
SEQ1903	CAATAGCTGGTATTGGTGGTTCAGGTAAAGCAGTTGCAGTTCAAGCAGCTATGGAGGGA
SEQ1904	TTTGAGCTGTTACGATTAAATAATCTAATTTCCGCAACTCCCTCC
SEQ1905	CAATAGCTGGTATTGGTGGTTCAGGTAAAGCAGTTGCAGTTCAAGCAGCTATGGAGGGA
SEQ1906	TTTGAGCTGTTACGATTAAATAATCTAATTTCCGCAACTCCCTCC
SEQ1907	
SEQ1908	TTTGAGCTATTACGATTAAATAATCTAATTTCCGCAACTCCCTCC
SEQ1909	CAATAGCTGGTATTGGTG
SEQ1910	TTTGAGCTGTTACGAT
SEQ1911	
SEQ1912	TTTGAGCTGTTACGATTAAATAATCTAATTTCCGCAAC
SEQ1913	CAATAGCTGGTATTGGTGGTTCAGGTAAAGCAGTTGCAGTTCAAGCAGCTATGGAGGGA
SEQ1914	TTTGAGCTATTACGATTAAATAATCTAATTTCCGCAACTCCCTCC

## Table 19: Comparative Sequences relating t SAF1680 (shikimate 5-dehydrogenase)

SEQ1901	GCAACTGCTTTACCTGAACCACCAATACCAGCTATTGTAATTATTTTATTTTAGCACT
SEQ1902	GCAACTGCTTTACCTGAACCACCAATACCAGCTATTGTAATTATTTTATTTTTAGCACT
SEQ1903	TTGCGGAAATTAGATTATTTAATCGTAACAGCTCAAATTACGATAAGGTCATTGACTTA
SEQ1904	GCAACTGCTTTACCTGAACCACCAATACCAGCTATTGTAATTATTTTATTTTTAGCACT
SEQ1905	TTGCGGAAATTAGATTATTTAATCGTAATAGCTCAAATTACGATAAGGTCATTGACTTA
SEQ1906	GCAACTGCTTTACCTGAACCACCAATACCAGCTATTGTAATTATTTTATTTTTAGCACT
	GCAACIGCI TACCIGAACACCACCAATACCAGCIAI IGTAATTATTI TATTI TAGCACT
SEQ1907	
SEQ1908	GCAACTGCTTTACCTGAACCACCAATACCAGCTATTGTAACTATTTT
SEQ1909	
SEQ1910	
SEQ1911	
SEQ1912	
	TTGCGG
SEQ1913	TTGCGG
SEQ1914	GCAACTGCTTTACCTGAACCACCAATACCAGCTATTGTAACTATTTTATTTTTAGCACT
SEQ1901	AAACCTTGAGCTGCTAAAGCTTTAAAACAACCAATGCCATCTGTCATATGGCCTACTAA
SEQ1902	AAACCTTGAGCTGCTAAAGCTTTAAAACAACCAATGCCATCTGTCATATGGCCTACTAA
SEQ1903	CAGATAAAATTAAAAAACAGTTTCAAATAAAGGTAGTCGTTGATTATCTAGAAAATAAG
SEQ1904	AAACCTTGAGCTGCTAAAGCTTTAAAACAACCAATGCCATCTGTCATATGGCCTACTAA
	CAGATAAAATTAAAAAACAGTTTCAAATAAAGGTAGTCGTTGATTATCTAGAAAATAAG
SEQ1905	
SEQ1906	AAACCTTGAGCTGCTAAAGCTTTAAAACAACCAATGCCATCTGTCATATGGCCTACTAA
SEQ1907	
SEQ1908	
SEQ1909	
SEQ1910	
SEQ1911	
SEQ1912	
SEQ1913	
SEQ1914	AAACCTTGAGCTGCTAAAGCTTTAAAACAACCAATGCCATCTGTCAT-TABCMARAT
SEQ1901	CGTCCGGTTCCACCTTGATTAACGATAGTATTTACAGCACCCACTAATTTAGCTTGAGG
SEQ1902	CGTCCGGTTCCACCTTGATTAACGATAGTATTTACAGCACCCACTAATTTAGCTTGAGG
	CAGCATTTAAAGACGCTATTAGAACTAGTCATTTTTATATTGATGCTACTAGTTTAGGA
SEQ1903	
SEQ1904	CGTCCGGTTCCACCTTGATTAACGATAGTATTTACAGCACCCACTAATTTAGCTTGAGG
SEQ1905	CAGCATTTAAAGACGCTATTAGAACTAGTCATTTTTATATTGATGCTACTAGTTTAGGA
SEQ1906	CGTCCGGTTCCACCTTGATTAACGATAGTATTTACAGCACCCACTAATTTAGCTTGAGG
SEQ1907	
SEQ1908	
SEQ1909	
SEQ1910	
SEQ1911	
SEQ1912	
SEQ1913	
SEQ1914	STNCSRATNGTSASHKMATDHYDRGNAS
SEQ1901	GATAAATCATCTAGCAAAGGGATAACACTCTGTTTAAATGGCATTGAAACATTAACACC
SEQ1902	GATAAATCATCTAGCAAAGGGATAACACTCTGTTTAAATGGCATTGAAACATTAACACC
SEQ1903	TGAGGCCATTAGATAATTATAGTTTAATTAACGATCCAGATATTTTAACACCGAATTTA
SEQ1904	GATAAATCATCTAGCAAAGGGATAACACTCTGTTTAAATGGCATTGAAACATTAACACC
SEQ1905	TGARGCCATTAGATAATTATAGTTTAATTAACGATCCAGAAATTTTAACACCCAATTTA
SEQ1906	GATAAATCATCTAGCAAAGGGATAACACTCTGTTTAAATGGCATTGAAACATTAACACC
SEQ1907	
SEQ1908	
SEQ1909	
SEQ1910	
SEQ1911	
SEQ1912	
SEQ1913	
SEQ1914	
-	

# Table 19: Comparative Sequences relating to SAF1680 (shikimat 5-dehydr genase)

_	
SEQ1901	CGAATACCCAATGCCCTGACACCTCGAACAGCTTCTGTTAATTTACCCTCTTCTACTTC
SEQ1902	CGAATACCCAATGCCCTGACACCTCGAACAGCTTCTGTTAATTTACCCTCTTCTACTTC
SEQ1903	TAGTTGTCGACTT
SEQ1904	CGAATACTCAATGCCCTGACACCTCGAACAGCTTCTGTTAATTTACCCTCTTCTACTTC
SEQ1905	TAGTTGTCGACTTGGTTTACAAGCCTAAAGAAACAGCATTGTTACGATTTGTTAGACAA
SEQ1906	CGAATACCCAATGCCCTGACACCTCGAACAGCTTCTGTTAATTTACCCTCTTCTACTTC
SEQ1907	
SEQ1908	
SEQ1909	
SEQ1910	
SEQ1911	
SEQ1912	
SE01913	
SE01914	
	·
SEQ1901	AATGTCAGATAGGCATAATTCATGTTTTTTTTTTTGAAAAGAGGTATTCCACATTAACGG
SE01902	AATGTCAGATAGGCATAATTCATGTTTTTTTTTTTGAAAAGAGGTATTCCACATTAACGG
SE01903	
SEQ1904	AATGTCAGATAGGCATAATTCATGTTTTTTTTTTTGAAAAGAGGTATTCCACATTAACGG
SEQ1905	ATGGAGTGAAACATGCTTATAATGGTCTAGGGATGCTGATTTATCAAGGAGCAGA
SEQ1906	AATGTCAGATAGGCATAATTCATGTTTTTTTTTTTGAAAAGAGGTATTCCACATTAACGG
SEQ1907	
SEQ1908	
SEQ1909	
SEQ1910	
SEQ1911	
SEQ1912	
SEQ1913	
SE01914	
SEQ1901	GATAGAGAGTGGCGTGCAGG-
SEQ1902	GATAGAGAGTGGCGTGCAGGA
SEQ1903	
SEQ1904	GATAGAGAGTGGCGTGCA
SEQ1905	
SEQ1906	GATAG
SEQ1907	
SEQ1908	
SEQ1909	w
SEQ1910	
SEQ1911	
SEQ1912	
SEQ1913	
SEQ1914	



#### Table 20: Comparative Sequences relating to SAG1723 (signal peptidase I)

# SEQ ID NO. 2002: SAG1680 FROM THE CJB110 GBS NONTYPEABLE STRAIN REVERSE COMPLEMENT

## Table 20: Comparative Sequences relating to SAG1723 (signal peptidas I)

SEQ2001	
SEQ2001 SEQ2002	
	TAAAGTTGACGGACACTCCATGGATCCAACTTTAGCTGACAAGGAACAGCTAGTAG
SEQ2003	TTGACGGACACTCCATGGATCCAACTTTAGCTGACAAGGAACAGCTAGTAG
SEQ2004 SEQ2005	AAGTTGACGGACACTCCATGGATCCAACTTTAGCTGACAAGGAACAGCTAGTAG
	TTGACGGACACTCCATGGATCCAACTTTAGCTGACAAGGAACAGCTAGTAG
SEQ2006	TTGACGGACACTCCATGGATCCAACTTTAGCTGACAAGGAACAGCTAGTAG
SEQ2007	TGGTAAAGTTGACGGACACTCCATGGATCCAACTTTAGCTGACAAGGAACAGCTAGTAG
SEQ2008	TTGACGGACACTCCATGGATCCAACTTTAGCTGACAAGGAACAGCTAGTAG
SEQ2009	TAAAGTTGACGGACACTCCATGGATCCAACTTTAGCTGACAAGGAACAGCTAGTAG
SEQ2010	AAAGTTGACGGACACTCCATGGATCCAACTTTAGCTGACAAGGAACAGCTAGTAG
SEQ2001	ATCGATTCGATATTGTAGTGGCTAACGAAGAAGAAGGCG
SEQ2002	TCTCAAACAAACAAAATCAATCGATTCGATATTGTAGTGGCTAACGAAGAAGAAGAGGCG
SEQ2002 SEQ2003	
SEQ2004	TCTCAAACAAACAAAATCAATCGATTCGATATTGTAGTGGCTAACGAAGAAGAAGGCG
SEQ2004 SEQ2005	TCTCAAACAAACAAAATCAATCGATTCGATATTGTAGTGGCTAACGAAGAAGAAGGCG
	TCTCAAACAAACAAAA—TAATCGATTCGATATTGTAGTGGCTAACGAAGAAGAAGGCG
SEQ2006	TCTCAAACAAACAAAATCAATCGATTCGATATTGTAGTGGCTAACGAAGAAGAAGGCG
SEQ2007	TCTCAAACAAACAAAATCAATCGATTCGATATTGTAGTGGCTAACGAAGAAGAAGGCG
SEQ2008	TCTCAAACAAACAAAATCAATCGATTCGATATTGTAGTGGCTAACGAAGAAGAAGGCG
SEQ2009	TCTCAAACAAACAAAATCAATCGATTCGATATTGTAGTGGCTAACGAAGAAGAAGGCG
SEQ2010	TCTCAAACAAACAAAATCAATCGATTCGATATTGTAGTGGCTAACGAAGAAGAAGGCG
SEQ2001	
SEQ2001	GCCAAAAGAAAAAATTGTTAAACGTGTCATTGGTATGCCAGGTGATGTCATCAAATATA
SEQ2003	GCCAAAAGAAAAATTGTTAAACGTGTCATTGGTATGCCAGGTGATGTCATCAAATATA
SEQ2004	GCCAAAAGAAAAAATTGTTAAACGTGTCATTGGTATGCCAGGTGATGTCATCAAATATA
SEQ2004 SEQ2005	GCCAAAAGAAAAAATTGTTAAACGTGTCATTGGTATGCCAGGTGATGTCATCAAATATA
-	GCCAAAAGAAAAAATTGTTAAACGTGTCATTGGTATGCCAGGTGATGTCATCAAATATA
SEQ2006	GCCAAAAGAAAAAATTGTTAAACGTGTCATTGGTATGCCAGGTGATGTCATCAAATATA
SEQ2007	GCCAAAAGAAAAATTGTTAAACGTGTCATTGGTATGCCAGGTGATGTCATCAAATATA
SEQ2008	GCCAAAAGAAAAATTGTTAAACGTGTCATTGGTATGCCAGGTGATGTCATCAAATATA
SEQ2009	GCCAAAAGAAAAAATTGTTAAACGTGTCATTGGTATGCCAGGTGATGTCATCAAATATA
SEQ2010	GCCAAAAGAAAAAATTGTTAAACGTGTCATTGGTATGCCAGGTGATGTCATCAAATATA
SEQ2001	<u> </u>
SEQ2002	AAAATGACACCTTAACTATTAACAATAAAAAAACAGAAGAACCTTACCTCAAGGAATATA
SEQ2003	AAAATGACACCTTAACTATTAACAATAAAAAAACAGAAGAACCTTACCTCAAGGAATATA
SEQ2003 SEQ2004	AAAATGACACCTTAACTATTAACAATAAAAAAACAGAAGAACCTTACCTCAAGGAATATA
	AAAATGACACCTTAACTATTAACAATAAAAAAACAGAAGAACCTTACCTCAAGGAATATA
SEQ2005	AAAATGACACCTTAACTATTAACAATAAAAAAACAGAAGAACCTTACCTCAAGGAATATA
SEQ2006	AAAATGACACCTTAACTATTAACAATAAAAAAACAGAAGAACCTTACCTCAAGGAATATA
SEQ2007	AAAATGACACCTTAACTATTAACAATAAAAAACAGAAGAACCTTACCTCAAGGAATATA
SEQ2008	AAAATGACACCTTAACTATTAACAATAAAAAAACAGAAGAACCTTACCTCAAGGAATATA
SEQ2009	AAAATGACACCTTAACTATTAACAATAAAAAAACAGAAGAACCTTACCTCAAGGAATATA
SEQ2010	AAAATGACACCTTAACTATTAACAATAAAAAAACAGAAGAACCTTACCTCAAGGAATATA
SEQ2001	C#A
SEQ2001 SEQ2002	CTAAATTATTT-AAAAAGGATAAATTACAGGAAAAATATTCGTATAACCCACTTTTCCAA
SEQ2002 SEQ2003	CTAAATTATTT-AAAAAGGATAAATTACAGGAAAAATATTCGTATAACCCACTTTTCCAA
SEQ2003 SEQ2004	CTAAATTATTT-AAAAAGGATAAATTACAGGAAAAATATTCGTATAACCCACTTTTCCAA
_	CTAAATTATTT-AAAAAGGATAAATTACAGGAAAAATATTCGTATAACCCACTTTTCCAA
SEQ2005	CTAAATTATTT-AAAAAGGATAAATTACAGGAAAAATATTCGTATAACCCACTTTTCCAA
SEQ2006	CTAAATTATTTTAAAAAGGATAAATTACAGGAAAAATATTCGTATAACCCACTTTTCCAA
SEQ2007	CTAAATTATTT-AAAAAGGATAAATTACAGGAAAAATATTCGTATAACCCACTTTTCCAA
SEQ2008	CTAAATTATTT-AAAAAGGATAAATTACAGGAAAAATATTCGTATAACCCACTTTTCCAA
SEQ2009	CTAAATTATTT-AAAAAGGATAAATTACAGGAAAAATATTCGTATAACCCACTTTTCCAA
SEQ2010	CTAAATTATTT-AAAAAGGATAAATTACAGGAAAAATATTCGTATAACCCACTTTTCCAA

# Table 20: C mparative Sequences relating to SAG1723 (signal peptidase I)

SEQ2001	GACCTAGCACAAAGCTCTACCGCTTTCACCACTGACAGCAATGGCAGCAGCGAATTTACT
SEQ2002	GACCTAGCACAAAGCTCTACCGCTTTCACTACTGACAGCAATTGGCAGCAGCGAATTTACT
SEQ2003	GACCTAGCACAAAGCTCTACCGCTTTCACCACTGACAGCAATGGCAGCAGCGAATTTACT
SEQ2004	GACCTAGCACAAAGCTCTACCGCTTTCACCACTGACAGCAATGGCAGCAGCGAATTTACT
SEQ2005	GACCTAGCACAAAGCTCTACCGCTTTCACCACTGACAGCAATGGCAGCAGCGAATTTACT
SEQ2006	GACCTAGCACAAAGCTCTACCGCTTTCACCACTGACAGCAATGGCAGCAGCGAATTTACT
-	
SEQ2007	GACCTAGCACAAAGCTCTACCGCTTTCACTACTGACAGCAATGGCAGCAGCGAATTTACC
SEQ2008	GACCTAGCACAAAGCTCTACCGCTTTCACCACTGACAGCAATGGCAGCAGCGAATTTACT
SEQ2009	GACCTAGCACAAAGCTCTACCGCTTTCACTACTGACAGCAATGGCAGCAGCGAATTTACT
SEQ2010	GACCTAGCACAAAGCTCTACCGCTTTCACCACTGACAGCAATGGCAGCAGCGAATTTACT
SEQ2001	CTGTCGTGCCTAAAGGCCACTACTATCTTGTTGGTGATGACCGAATTGTCTCTAAAGAT
SEQ2002	CTGTCGTGCCTAAAGGCCACTATTATCTTGTTGGTGATGACCGAATTGTCTCTAAAGAT
SEQ2003	CTGTCGTGCCTAAAGGCCACTACTATCTTGTTGGTGATGACCGAATTGTCTCTAAAGAT
SEQ2004	CTGTCGTGCCTAAAGGCCACTACTATCTTGTTGGTGATGACCGAATTGTCTCTAAAGAT
SEQ2005	CTGTCGTGCCTAAAGGCCACTACTATCTTGTTGGTGATGACCGA
SEQ2006	oldiddiwddodd Holaidi Holaidd
SEQ2007	CTGTCGTGCCTAAAGGCCACTACTATCTTGTTGGTGATGACCGAATTGTCTCTAAAGAT
SEQ2008	CTGTCGTGCCTAAAGGCCACTACTATCTTGTTGGTGATGACCGAATTGTCTCTAAAGAT
SEQ2008 SEQ2009	CTGTCGTGCCTAAAGGCCACTATTATCTTGTTGGTGATGACCGAATTGTCTCTAAAGAT
SEQ2010	CTGTCGTGCCTAAAGGCCACTACTATCTTGTTGGTGATGACCGAATTGTCTCTAAAGAT
SEQ2001	GTCGTGCCGTCGGTTCCTTCAAAA
SEQ2002	GTCGTGCCGTCCGTTCAAAAATCAACAATTGTGGGAG
SEQ2003	
3EQ2003	GTCGTGCCGTCGGTCCCTTCAAAAAATCAACGATTGTGGGAGAGGT
SEQ2003 SEQ2004	GTCGTGCCGTCGGTCCCTTCAAAAAATCAACGATTGTGGGAGAGGTGTCGTGCCGTCGGT
	GTCGTGCCGTCGGTCCCTTCAAAAAATCAACGATTGTGGGAGAGGTGTCGTGCCGTCGGT
SEQ2004 SEQ2005	GTCGTGCCGTCGCTTCAAAAAATCAACGATTGTGGGAGAGGTGTCGTGCCGTCGGT
SEQ2004 SEQ2005 SEQ2006	GTCGTGCCGTCGGT
SEQ2004 SEQ2005 SEQ2006 SEQ2007	GTCGTGCCGTCGGT
SEQ2004 SEQ2005 SEQ2006 SEQ2007 SEQ2008	GTCGTGCCGTCGGT
SEQ2004 SEQ2005 SEQ2006 SEQ2007 SEQ2008 SEQ2009	GTCGTGCCGTCGGT
SEQ2004 SEQ2005 SEQ2006 SEQ2007 SEQ2008	GTCGTGCCGTCGGT
SEQ2004 SEQ2005 SEQ2006 SEQ2007 SEQ2008 SEQ2009	GTCGTGCCGTCGGT
SEQ2004 SEQ2005 SEQ2006 SEQ2007 SEQ2008 SEQ2009 SEQ2010	GTCGTGCCGTCGGT
SEQ2004 SEQ2005 SEQ2006 SEQ2007 SEQ2008 SEQ2009 SEQ2010	GTCGTGCCGTCGGT
SEQ2004 SEQ2005 SEQ2006 SEQ2007 SEQ2008 SEQ2009 SEQ2010 SEQ2001	GTCGTGCCGTCGGT
SEQ2004 SEQ2005 SEQ2006 SEQ2007 SEQ2008 SEQ2009 SEQ2010 SEQ2001 SEQ2001	GTCGTGCCGTCGGT
SEQ2004 SEQ2005 SEQ2006 SEQ2007 SEQ2008 SEQ2009 SEQ2010 SEQ2001 SEQ2001 SEQ2002 SEQ2003 SEQ2004 SEQ2005	GTCGTGCCGTCGGT
SEQ2004 SEQ2005 SEQ2006 SEQ2007 SEQ2008 SEQ2009 SEQ2010 SEQ2001 SEQ2001 SEQ2002 SEQ2003 SEQ2004 SEQ2005 SEQ2006	GTCGTGCCGTCGGT
SEQ2004 SEQ2005 SEQ2006 SEQ2007 SEQ2008 SEQ2009 SEQ2010 SEQ2001 SEQ2002 SEQ2003 SEQ2004 SEQ2005 SEQ2006 SEQ2007	GTCGTGCCGTCGGT
SEQ2004 SEQ2005 SEQ2006 SEQ2007 SEQ2008 SEQ2009 SEQ2010 SEQ2001 SEQ2002 SEQ2003 SEQ2004 SEQ2005 SEQ2006 SEQ2007 SEQ2008	GTCGTGCCGTCGGT
SEQ2004 SEQ2005 SEQ2006 SEQ2007 SEQ2008 SEQ2009 SEQ2010 SEQ2001 SEQ2002 SEQ2003 SEQ2004 SEQ2005 SEQ2006 SEQ2007	GTCGTGCCGTCGGT



### 21: Comparative Sequences relating to SAG0079 (adenylate kinase)

SET ID NO. 2101: SAG0079 FROM THE 2603V/R GBS TYPE V STRAIN
AATCTTTTAATTATGGGTTTGCCTGGTGCTGGTAAAGGTACTCAAGCAGCTAAGATCGTTGAAGAATTTGGTGTTGCTCACATCTCA
ACAGGGGATATGTTCCGCGCCGCAATGGCTAAAACGGAATCGAAACGGACGTTAGCTAAAAGGTTATATTGATAAAGGTGAATTGGTT
CCTGATGAAGAACAAACGGGATTGTAAAAAGAGCGCTTAGCTGAGGATGATATCGCAGAAAAAAGGTTTTTTACTTGATGGATATCCA
CGTACTATTGAACAAGCACACGCCTTAGATGCTACGCTTGAAGAACTAGGACTACGCTTAGATGGTGTTATTAATATTAAAGTGGAT
CCATCATGTCTTATAGAGCGTTTGAGTGGTCGTATTATCAATCGTAAAACTGTGAAACTTTCCACAAAGTGTTCAACCCACCAGTA
GATTATAAAAGAAGAAGATTACTATCAACGTGAAGATGATAAGCCTGAAACTGTCAAACGTCGCTTGGACGTTAATATTGCTCAAGGA
GAACCTATTCTTGAACACTATCGTAAGCTTTGGTCTTTACAGATATTTGAAGGTAATCAAGAAATAACAGAAGTTTTTTGCAGATGTT
GAAAAAGCGTTG

SEQ ID NO. 2102: SAG0079 FROM THE 090 GBS TYPE IA STRAIN REVERSE COMPLEMENT
AATCTTTTAATTATGGGTTTGCCTGGTGCTGGTAAAGGTACTCAAGCAGCTAAGATCGTTGAAGAATTTGGTGTTGCTCACATCTCA
ACAGGGGATATGTTCCGCGCCCGCAATGGCTAATCAAACCGAAATGGGACGTTTAGCTAAAAGGTTATATTGATAAAGGTGAATTGGTT
CCTGATGAAGTAACAAACGGGATTGTAAAAGAGCGCTTAGCTGAGGATGATATCCACAGAAAAAAGGTTTTTTACTTGATGGATATCCA
CGTACTATTGAACAAGCACACCCCTTAGATGCTACGCTTGAAGAACTAGGACTTACACAGAAGTGTTCAACCCACAGTA
CATCATGTCTTATAGAGCGTTTGAGTGGTCGTATTATCAATCGTAAAACTGTCAAACCTTCCACAAAGTGTTCAACCCACCAGTA
GATTATAAAGAAGAAGATTACTATCAACGTGAAGATGATAAACTGTCAAACGTCGCTTGGACGTTAATATTGCTCAAGGA
GAACCTATTCTTGAACACTATCGTAAGCTTGGTCTTGTTACAGATATTGAAGGTAATCAAGAAATAACAGAAGTTTTTGCAGATGTT
GAAAAAGCGTTGCTAGGACTCAAA

SEQ ID NO. 2104: SAG0079 FROM THE 18RS21 GBS TYPE II STRAIN REVERSE COMPLEMENT
AATCTTTTAACCACGGGTTCGCCTGGTGCTGGTAAAGGTACTCAAGCAGCTAAGATCGTTGAAGAATTTGGTGTTGCTCACATCTCA
ACAGGGGATATGTTCCGCGCCCGCAATGGCTAATCAAACCGAAATGGGACGTTTAGCTAAAAGGTTATATTGATAAAGGTGAATTGGTT
CCTGATGAAGAACAAACGGGATTGTAAAAGAGCGCTTAGGTGAGAACTACGCAGAAAAAGGTTTTTTACTTGATGGATATCCA
CGTACTATTGAACAAGCACACGCCTTAGATGCTACGCTTGAAGAACTAGGACTACGCTTAGATGGTGTTATTAATATTAAAGTGGAT
CCATCATGTCTTATAAGAGCGTTTGAGTGGTCGTATTATCAATCGTAAAACTGGTGAAACTTTCCACAAAGTGTTCAACCCACCAGTA
GATTATAAAGAAGAAGATTACTATCAACGTGAAGATGATAAGCCTGAAACTGTCAAAACGTCGCTTGGACGTTAATATTGCTCAAGGA
GAACCTATTCTTGAACACATATCGTAAGCTTGGTCTTTGTTACAGATATTGAAGGTAATCAAGAAATAACAGAAGTTTTTGCAGATGTT
GAAAAAAGCGTTGCTAGGA

SEQ ID NO. 2106: SAG0079 FROM THE A909 GBS TYPE Ia STRAIN REVERSE COMPLEMENT
AATCTTTTAATTATGGGTTTGCCTGGTGCTGGTAAAGGTACTCAAGCAGCTAAGATCGTTGAAGAATTTGGTGTTGCTCACATCTCA
ACAGGGGATATGTTCCGCGCCGCAATGGCTAATCAAACCGAAATGGGACGTTTAGCTAAAAGGTTATATTGATAAAGGTGAATTGGTT
CCTGATGAAGAACAAACGGGATTGTAAAAGAGCGCTTAGCTGAAGAATTATCGCAGAAAAAGGTTTTTTACTTGATGGATATCCA
CGTACTATTGAACAAGCACACGCCTTAGATGCTACGCTTGAAGAACTAGGACTACGCTTAGATGGTGTTATTAAAATTAAAGTGGAT
CCATCATGTCTTATAGAGCGTTTGAGTGGTCGTATTATCAATCGTAAAACTGGTGAAACTTTCCACAAAGTGTTCAACCCACCAGTA
GATTATAAAGAAGAAGATTACTATCAACGTGAAGATGATAAGCCTGAAACTGTCAAACGTCGCTTGGACGTTAATATTGCTCAAGGA
GAATCTATTCTTGAACACTATCGAAAGCTTGGTCTTGTTACAGATATTGAAGGTAA

SEQ ID NO. 2107: SAG0079 FROM THE CJB110 GBS NONTYPEABLE STRAIN REVERSE COMPLEMENT AATCTTTAACCACGGGTTTGCTTGGTGGTGAAAGGTACTCAAGCAGCTAAGATCGTTGAAGAATTTGGTGTTGCTCACATCTCA ACAGGGGATATGTTCCGCGCCGCAATGGCTAAACCGAAATGGGACGTTTAGCTAAAAGTTATATTGATAAAGGTGAATTCCA CCTGATGAAGAACAAACGGGATTGTAAAAGGGGATTGTAAAAGGTGATATCCA CGTACTATTGAACAACACGCCTTAGATGCTACGACTAACACTAGGACTACGCTTAGATGGTTTATTAAATATTAAAGTGGAT CCATCATGTCTTATAGAGCGGTTTGATGGTCGTATTATCAATCGTAAAACTGGTGAAACTTTCCACAAAGTGTTCAACCCACCAGTA GATTATAAAGAAGAAGAAGATTACTATCAACGTGAAGATAAAGCTGTCAAACGTCGCTTGGACGTTAATATTGCTCAAGGA GAACCTATTCTTGAACACTATAG

### Time 21: Comparative Sequences relating to SAG0079 (adenylate kinase)

SEQ ID NO. 2109: SAG0079 FROM THE H36b GBS TRYP Ib STRAIN REVERSE COMPLEMENT CAGGGGATATGTTCCGCGCCGCAATGGCTAATCAAACCGAAATGGGACGTTTAGCTAAAAGTTATATTGATAAAGGTGAATTGGTTC CTGATGAAGATAACAAACGGGATTGTAAAAGAGGCCTTAGCTGAGGATGATATCGCAGAAAAAAGGTTTTTTACTTGATGGATATCCAC GTACTATTGAACAAGCACACGCCTTAGATGCTTGAAGAACTAGGACTACGCTTAGATGGTGTTATTAATATTAAAGTGGATC CATCATGTCTTATAGAGCGTTTGAGTGGTCGTATTATCAATCGTAAAACTGTGAAACTTTCCACAAAGTGTTCAACCCACCAGTAGATTATAAAAGAAGAAGAAGTTACTATCAACGTGAAAACTGTCAAACGTCGCTTGGACGTTAATATTTCCTCAAGGAGAATTATTATTAGATGTTTGAACGTTATCATTCTTGAACACTATCGTAAGCTTTGTACAGAATTTCAAGGAGAATAACAGAAGTTTTTTGCAGATGTTGAAAAGCGTTGCT

SEQ ID NO. 2112: SAG0079 FROM THE M781 GBS TYPE III STRAIN REVERSE COMPLEMENT
AATCTTTTAATTACGGGTTTGCCTGGTGCTGGTAAAGGTACTCAAGCAGCTAAGATTGTTGAAGAATTTGGTGTTGCTCACATCTCA
ACAGGGGATATGTTCCGCGCCCGCAATGGCTAATCAAACCCAAATGGGACGTTTAGCTAAAAGGTTATATTGATAAAGGTGAATTGGTT
CCTGATGAAGTAACAAACGGGATTGTAAAAGAGCGCTTAGCTGAAGAATTATCGCAGAAAAAAGGTTTTTTACTTGATGGATATCCA
CGTACTATTGAGCAAGCACACGCCTTAGATGCTACGCTTGAAGAACTAGGACTACGCTTAGATGGTGTTATTAAATATTAAAGTGGAT
CCAACATGCCTTATAGAGCGTTTGAGTGGCCGTATTATCAATCGTAAAACTGGTGAAACTTTCCACAAAGTGTTCAACCCCACCAGTA
GATTATAAAGAAGAAGATTACTATCAACGTGAAGATGATAAGCCTGAAACTGTCAAACGTCGCTTGGACGTTAATATTGCTCAA

SEQ2101	ATCTTTAATTATGGGTTTGCCTGGTGCTGGTAAAGGTACTCAAGCAGCTAAGATCGTT
SEQ2102	ATCTTTAATTATGGGTTTGCCTGGTGCTGGTAAAGGTACTCAAGCAGCTAAGATCGTT
SEQ2103	tggtaaagggactcaagcagctaagattgtt
SEQ2104	ATCTTTTAACCACGGGTTCGCCTGGTGCTGGTAAAGGTACTCAAGCAGCTAAGATCGTT
SEQ2105	ATCTTTAATTATGGGTTŢGCCTGGTGCTGGTAAAGGTACTCAAGCAGCTAAGATCGTT
SEQ2106	ATCTTTTAATTATGGGTTTGCCTGGTGCTGGTAAAGGTACTCAAGCAGCTAAGATCGTT
SEQ2107	ATCTTTTAACCACGGGTTTGCTTGGTGCTGGTAAAGGTACTCAAGCAGCTAAGATCGTT
SEQ2108	ATCTTTAATTATGGGTTTGCCTGGTGCTGGTAAAGGTACTCAAGCAGCTAAGATTGTT
SEQ2109	
SEQ2110	ATCTTTTAATTATGGGTTTGCCTGGTGCTGGTAAAGGTACTCAAGCAGCTAAGATCGTT
SEQ2111	CTTTTAATTATGGGTTTGCCTGGTGCTGGTAAAGGTACTCAAGCAGCTAAGATTGTT
SEQ2112	ATCTTTAATTACGGGTTTGCCTGGTGCTGGTAAAGGTACTCAAGCAGCTAAGATTGTT
SEQ2101	AAGAATTTGGTGTTGCTCACATCTCAACAGGGGATATGTTCCGCGCCGCAATGGCTAAT
SEQ2102	AAGAATTTGGTGTTGCTCACATCTCAACAGGGGATATGTTCCGCGCCGCAATGGCTAAT
SEQ2103	AAGAATTTGGTGTTGCGCACATCTCAACAGGGGATATGTTCCGCGCCGCAATGGCTAAT
SEQ2104	AAGAATTTGGTGTTGCTCACATCTCAACAGGGGATATGTTCCGCGCCGCAATGGCTAAT
SEQ2105	AAGAATTTGGTGTTGCTCACATCTCAACAGGGGATATGTTCCGCGCCGCAATGGCTAAT
SEQ2106	AAGAATTTGGTGTTGCTCACATCTCAACAGGGGATATGTTCCGCGCCGCAATGGCTAAT
SEQ2107	AAGAATTTGGTGTTGCTCACATCTCAACAGGGGATATGTTCCGCGCCGCAATGGCTAAT
SEQ2108	AAGAATTTGGTGTTGCTCACATCTCAACAGGGGATATGTTCCGCGCCGCAATGGCTAAT
SEQ2109	
SEQ2110	AAGAATTTGGTGTTGCTCACATCTCAACAGGGGATATGTTCCGCGCCGCAATGGCTAAT

# Table 21: Comparative Sequences relating to SAG0079 (adenylate kinase)

SECTION SEQUENCE	AAGAATTTGGTGTTGCTCACATCTCAACAGGGGATATGTTCCGCGCCCAATGGCTAAT AAGAATTTGGTGTTGCTCACATCTCAACAGGGGATATGTTCCGCGCCCCAATGGCTAAT
SEQ2101	Caaaccgaaatgggacgtttagctaaaagttatatttgataaaggtgaattggttcctgat Caaaccgaaatgggacgtttagctaaaagttatattgataaaggtgaattggttcctgat
SEQ2102	CAAACCGAAATGGGACGTTTAGCTAAAAGTTATATTGATAAAGGTGAATTGGTTCCTGAT CAAACCGAAATGGGACGTTTAGCTAAAAGTTATATTGATAAAGGTGAATTGGTTCCTGAT
SEQ2103	CAAACCGAAATGGGACGTTTAGCTAAAAGTTATATTGATAAAGGTGAATTGGTTCCTGAT
SEQ2104	CAAACCGAAATGGGACGTTTAGCTAAAAGTTATATTGATAAAGGTGAATTGGTTCCTGAT
SEQ2105	CAAACCGAAATGGGACGTTTAGCTAAAAGTTATATTGATAAAGGTGAATTGGTTCCTGAT
SEQ2106	CAAACCGAAATGGGACGTTTAGCTAAAAGTTATATTGATAAAGGTGAATTGGTTCCTGAT
SEQ2107	CAAACCGAAATGGGACGTTTAGCTAAAAGTTATATTGATAAAGGTGAATTGGTTCCTGAT
SEQ2108 SEO2109	CAAACCCAAATGGGACGTTTAGCTAAAAGTTATATTGATAAAGGTGAATTGGTTCCTGAT
SEQ2109 SEQ2110	CAAACCGAAATGGGACGTTTAGCTAAAAGTTATATTGATAAAGGTGAATTGGTTCCTGAT
-	CAAACCCAAATGGGACGTTTAGCTAAAAGTTATATTGATAAAGGTGAATTGGTTCCTGAT
SEQ2111 SEQ2112	CAAACCCAAATGGGACGTTTAGCTAAAAGTTATATTGATAAAGGTGAATTGGTTCCTGAT
D-12	
SEQ2101	AAGTAACAAACGGGATTGTAAAAGAGCGCTTAGCTGAGGATGATATCGCAGAAAAAGGT
SEQ2102	AAGTAACAAACGGGATTGTAAAAGAGCGCTTAGCTGAGGATGATATCGCAGAAAAAGGT
SEQ2103	AAGTAACAAACGGGATTGTAAAAGAGCGCTTAGCTGAGGATGATATCGCAGAAAAAGGT
SEQ2104	AAGTAACAAACGGGATTGTAAAAGAGCGCTTAGCTGAGGATGATATCGCAGAAAAAGGT
SEQ2105	AAGTAACAAACGGGATTGTAAAAGAGCGCTTAGCTGAGGATGATATCGCAGAAAAAGGT
SEQ2106	. AAGTAACAAACGGGATTGTAAAAGAGCGCTTAGCTGAGGATGATATCGCAGAAAAAGGT
SEQ2107	AAGTAACAAACGGGATTGTAAAAGAGCGCTTAGCTGAGGATGATATCGCAGAAAAAGGT
SEQ2108	AAGTAACAAACGGGATTGTAAAAGAGCGCTTAGCTGAGGATGATATCGCAGAAAAAGGT
SEQ2109	AAGTAACAAACGGGATTGTAAAAGAGCGCTTAGCTGAGGATGATATCGCAGAAAAAGGT
SEQ2110	AAGTAACAAACGGGATTGTAAAAGAGCGCTTAGCTGAGGATGATATCGCAGAAAAAGGT
SEQ2111	AAGTAACAAACGGGATTGTAAAAGAGCGCTTAGCTGAGGATGATATCGCAGAAAAAGGT
SEQ2112	AAGTAACAAACGGGATTGTAAAAGAGCGCTTAGCTGAGGATGATATCGCAGAAAAAGGT
SEQ2101	TTTTTACTTGATGGATATCCACGTACTATTGAACAAGCACACGCCTTAGATGCTACGCTT
SEQ2102	TTTTTACTTGATGGATATCCACGTACTATTGAACAAGCACACGCCTTAGATGCTACGCTT
SEQ2103	TTTTTACTTGATGGGTATCCACGTACTATTGAACAAGCACACGCCTTAGATGCTACGCTT
SEQ2104	TTTTTACTTGATGGATATCCACGTACTATTGAACAAGCACACGCCTTAGATGCTACGCTT
SEQ2105	TTTTTACTTGATGGATATCCACGTACTATTGAACAAGCACACGCCTTAGATGCTACGCTT
SEQ2106	TTTTTACTTGATGGATATCCACGTACTATTGAACAAGCACACGCCTTAGATGCTACGCTT
SEQ2107	TTTTTACTTGATGGATATCCACGTACTATTGAACAAGCACACGCCTTAGATGCTACGCTT
SEQ2108	TTTTTACTTGATGGATATCCACGTACTATTGAGCAAGCACACGCCTTAGATGCTACGCTT
SEQ2109	TTTTTACTTGATGGATATCCACGTACTATTGAACAAGCACACGCCTTAGATGCTACGCTT
SEQ2110	TTTTTACTTGATGGATATCCACGTACTATTGAACAAGCACACGCCTTAGATGCTACGCTT
SEQ2111	TTTTTACTTGATGGATATCCACGTACTATTGAGCAAGCACACGCCTTAGATGCTACGCTT
SEQ2112	TTTTTACTTGATGGATATCCACGTACTATTGAGCAAGCACACGCCTTAGATGCTACGCTT
SEQ2101	GAAGAACTAGGACTACGCTTAGATGGTGTTATTAATATTAAAGTGGATCCATCATGTCTT
SEQ2102	GAAGAACTAGGACTACGCTTAGATGGTGTTATTAATATTAAAGTGGATCCATCATGTCTT
SEQ2103	GAAGAACTAGGACTACGCTTAGATGGTGTTATTAATATTAAAGTGGATCCATCATGTCTT
SEQ2104	GAAGAACTAGGACTACGCTTAGATGGTGTTATTAATATTAAAGTGGATCCATCATGTCTT
SEQ2105	GAAGAACTAGGACTACGCTTAGATGGTGTTATTAATATTAAAGTGGATCCATCATGTCTT
SEQ2106	GAAGAACTAGGACTACGCTTAGATGGTGTTATTAATATTAAAGTGGATCCATCATGTCTT
SEQ2107	GAAGAACTAGGACTACGCTTAGATGGTGTTATTAATATTAAAGTGGATCCATCATGTCTT
\$EQ2108	GAAGAACTAGGACTACGCTTAGATGGTGTTATTAATATTAAAGTGGATCCAACATGCCTT
SEQ2109	GAAGAACTAGGACTACGCTTAGATGGTGTTATTAATATTAAAGTGGATCCATCATGTCTT
SEQ2110	GAAGAACTAGGACTACGCTTAGATGGTGTTATTAATATTAAAGTGGATCCATCATGTCTT
SEQ2111	GAAGAACTAGGACTACGCTTAGATGGTGTTATTAATATTAAAGTGGATCCAACATGCCTT
SEQ2112	GAAGAACTAGGACTACGCTTAGATGGTGTTATTAATATTAAAGTGGATCCAACATGCCTT
SEQ2101	ATAGAGCGTTTGAGTGGTCGTATTATCAATCGTAAAACTGGTGAAACTTTCCACAAAGTG
SEQ2102	ATAGAGCGTTTGAGTGGTCGTATTATCAATCGTAAAACTGGTGAAACTTTCCACAAAGTG
SEQ2103	ATAGAGCGTTTGAGTGGTCGTATTATCAATCGTAAAACTGGTGAAACTTTCCACAAAGTG
SEQ2104	ATAGAGCGTTTGAGTGGTCGTATTATCAATCGTAAAACTGGTGAAACTTTCCACAAAGTG
SEQ2105	ATAGAGCGTTTGAGTGGTCGTATTATCAATCGTAAAACTGGTGAAACTTTCCACAAAGTG
SEQ2105	ATAGAGCGTTTGAGTGGTCGTATTATCAATCGTAAAACTGGTGAAACTTTCCACAAAGTG
SEQ2103	ATAGAGCGTTTGAGTGGTCGTATTATCAATCGTAAAACTGGTGAAACTTTCCACAAAGTG
SEQ2107	ATAGAGCGTTTGAGTGGCCGTATTATCAATCGTAAAACTGGTGAAACTTTCCACAAAGTG
SEQ2108	ATAGAGCGTTTGAGTGGTCGTATTATCAATCGTAAAACTGGTGAAACTTTCCACAAAGTG
	ATAGAGCGTTTGAGTGGTCGTATTATCAATCGTAAAACTGGTGAAACTTTCCACAAAGTG
SEQ2110	ATAGAGCGTTTGAGTGGTCGTATTATCAATCGTAAAACTGGTGAAACTTTCCACAAAGTG ATAGAGCGTTTGAGTGGCCGTATTATCAATCGTAAAACTGGTGAAACTTTCCACAAAGTG
SEQ2111	ATAGAGCGTTTGAGTGGCCGTATTATCAATCGTAAAACTGGTGAAACTTTCCACAAAGTG ATAGAGCGTTTGAGTGGCCGTATTATCAATCGTAAAACTGGTGAAACTTTCCACAAAGTG
SEQ2112	ATAGAGCGTTTGAGTGGCCGTATTATCAATCGTAAAACTGGTGAAACTTTCCACAAAGTG

## Table 21: Comparative Sequences relating to SAG0079 (adenylate kinase)

SECTION	TTCAACCCACCAGTAGATTATAAAGAAGAAGATTACTATCAACGTGAAGATGATAAGCCT
SEQ2102	TTCAACCCACCAGTAGATTATAAAGAAGAAGATTACTATCAACGTGAAGATGATAAGCCT
SEQ2103	TTCAACCCACCAGTAGATTATAAAGAAGAAGATTACTATCAACGTGAAGATGATAAGCCT
SEQ2104	TTCAACCCACCAGTAGATTATAAAGAAGAAGATTACTATCAACGTGAAGATGATAAGCCT
SEQ2105	TTCAACCCACCAGTAGATTATAAAGAAGAAGATTACTATCAACGTGAAGATGATAAGCCT
SEQ2106	TTCAACCCACCAGTAGATTATAAAGAAGAAGATTACTATCAACGTGAAGATGATAAGCCT
SEQ2107	TTCAACCCACCAGTAGATTATAAAGAAGAAGATTACTATCAACGTGAAGATGATAAGCCT
SEQ2108	TTCAACCCACCAGTAGATTATAAAGAAGAAGATTACTATCAACGTGAAGATGATAAGCCT
SEQ2109	TTCAACCCACCAGTAGATTATAAAGAAGAAGATTACTATCAACGTGAAGATGATAAGCCT
SEQ2110	TTCAACCCACCAGTAGATTATAAAGAAGAAGATTACTATCAACGTGAAGATGATAAGCCT
SEQ2111	TTCAACCCACCAGTAGATTATAAAGAAGAAGATTACTATCAACGTGAAGATGATAAGCCT
SEQ2112	TTCAACCCACCAGTAGATTATAAAGAAGAAGATTACTATCAACGTGAAGATGATAAGCCT
SEQ2101	GAAACTGTCAAACGTCGCTTGGACGTTAATATTGCTCAAGGAGAACCTATTCTTGAACAC
SEQ2102	GAAACTGTCAAACGTCGCTTGGACGTTAATATTGCTCAAGGAGAACCTATTCTTGAACAC
SEQ2103	GAAACTGTCAAACGTCGCTTGGACGTTCATATTGCTCAAGGAGAACCTATTCTTGAACAC
SEQ2104	GAAACTGTCAAACGTCGCTTGGACGTTAATATTGCTCAAGGAGAACCTATTCTTGAACAC
SEQ2105	GAAACTGTCAAACGTCGCTTGGACGTTAATATTGCTCAAGGAGAACCTATTCTTGAACAC
SEQ2106	GAAACTGTCAAACGTCGCTTGGACGTTAATATTGCTCAAGGAGAATCTATTCTTGAACAC
SEQ2107	GAAACTGTCAAACGTCGCTTGGACGTTAATATTGCTCAAGGAGAACCTATTCTTGAACAC
SEQ2108	GAAACTGTCAAACGTCGCTTGGACGTTAATATTGCTCAAGGAGAACCTATTCTTGAACAC
SEQ2109	GAAACTGTCAAACGTCGCTTGGACGTTAATATTGCTCAAGGAGAATCTATTCTTGAACAC
SEQ2110	GAAACTGTTAAACGTCGCTTGGACGTTAATATTGCTCAAGGAGAACCTATTCTTGAACAC
SEQ2111	GAAACTGTCAAACGTCGCTTGGACGTTAATATTGCTCAAGGAGAACCTATTCTTGAACAC
SEQ2112	GAAACTGTCAAACGTCGCTTGGACGTTAATATTGCTCAATABCMARATVSTNCSRAT
SEQ2101	ATCGTAAGCTTGGTCTTGTTACAGATATTGAAGGTAATCAAGAAATAACAGAAGTTTTT
SEQ2102	ATCGTAAGCTTGGTCTTGTTACAGATATTGAAGGTAATCAAGAAATAACAGAAGTTTTT
SEQ2103	ATAGTAAGCTTGGCCTTGTTACAGATATTGAAGGTAATCAAGAAATAA
SEQ2104	ATCGTAAGCTTGGTCTTGTTACAGATATTGAAGGTAATCAAGAAATAACAGAAGTTTTT
SEQ2105	ATCGTAAGCTTGGTCTTGTTACAGATATTGAAGGTAATCAAGAAATAACAGAAGTTTTT
SEQ2106	ATCGAAAGCTTGGTCTTGTTACAGATATTGAAGGTAA
SEQ2107	ATAG
SEQ2108	ATCGTAAGCTTGGTCTTGTTACAGATATTGAAGGTAATCAAGAAATAACAGAAGTTTTT
SEQ2109	ATCGTAAGCTTGGTCTTGTTACAGATATTGAAGGTAATCAAGAAATAACAGAAGTTTTT
SEQ2110	ATAAAAAGCTTGGTCTTGTTACAGATATTGAAGGTAATCA
SEQ2111	ATCGTAAGCTTGGTCTTGTTACAGATATTGAAGGTAATCAAGAAATAACAGAAGTTTTT
SEQ2112	GTSAGADNYATKNAS
SEQ2101	CAGATGTTGAAAAAGCGTTG
SEQ2102	CAGATGTTGAAAAAGCGTTGCTAGAACTCAAA
SEQ2103	
SEQ2104	CAGATGTTGAAAAAGCGTTGCTAGAA
SEQ2105	CAGATGTTGAAAAAGCGTTG
SEQ2106	
SEQ2107	
SEQ2108	CAGATGTTGAAAAAGCGTTGCTAG
SEQ2109	
	CAGATGTTGAAAAAGCGTTGCT
SEQ2110	CAGATGTTGAAAAAGCGTTGCT



## Table 21: C mparative Sequences relating t SAG0079 (adenylate kinase)

#### >SEO ID NO 2150:090 frame: 1

NLLIMGLPGAGKGTQAAKIVEEFGVAHISTGDMFRAAMANQTEMGRLAKSYIDKGELVPD EVTNGIVKERLAEDDIAEKGFLLDGYPRTIEQAHALDATLEELGLRLDGVINIKVDPSCL IERLSGRIINRKTGETFHKVFNPPVDYKEEDYYQREDDKPETVKRRLDVNIAQGEPILEH YRKLGLVTDIEGNQEITEVFADVEKALLELK

#### >SEQ ID NO 2151:114\_1169NT frame: 2

GKGTQAAKIVEEFGVAHISTGDMFRAAMANQTEMGRLAKSYIDKGELVPDQVTNGIVKER LAEDDIAEKGFLLDGYPRTIEQAHALDATLEELGLRLDGVINIKVDPSCLIERLSGRIIN RKTGETFHKVFNPPVDYKEEDYYQREDDKPETVKRRLDVHIAQGEPILEHYSKLGLVTDI EGNOEI

#### >SEQ ID NO 2152: 114\_18RS21 frame: 1

NILTTGSPGAGKGTQAAKIVEEFGVAHISTGDMFRAAMANQTEMGRLAKSYIDKGELVPD EVTNGIVKERLAEDDIAEKGFLLDGYPRTIEQAHALDATLEELGLRLDGVINIKVDPSCL IERLSGRIINRKTGETFHKVFNPPVDYKEEDYYQREDDKPETVKRRLDVNIAQGEPILEH YRKLGLVTDIEGNQEITEVFADVEKALLE

#### >SEQ ID NO 2153: 114 2603 frame: 1

NILIMGIPGAGKGTQAAKIVEEFGVAHISTGDMFRAAMANQTEMGRLAKSYIDKGELVPD EVTNGIVKERLAEDDIAEKGFLLDGYPRTIEQAHALDATLEELGLRLDGVINIKVDPSCL IERLSGRIINRKTGETFHKVFNPPVDYKEEDYYQREDDKPETVKRRLDVNIAQGEPILEH YRKLGLVTDIEGNQEITEVFADVEKAL

#### >SEQ ID NO 2154: 114 A909 frame: 1

NLLIMGLPGAGKGTQAAKIVEEFGVAHISTGDMFRAAMANQTEMGRLAKSYIDKGELVPD EVTNGIVKERLAEDDIAEKGFLLDGYPRTIEQAHALDATLEELGLRLDGVINIKVDPSCL IERLSGRIINRKTGETFHKVFNPPVDYKEEDYYQREDDKPETVKRRLDVNIAQGESILEH YRKLGLVTDIEG

#### >SEQ ID NO 2155:114 A909 frame: 1

NILIMGLPGAGKGTQAAKIVEEFGVAHISTGDMFRAAMANQTEMGRLAKSYIDKGELVPD EVTNGIVKERLAEDDIAEKGFLLDGYPRTIEQAHALDATLEELGLRLDGVINIKVDPSCL IERLSGRIINRKTGETFHKVFNPPVDYKEEDYYQREDDKPETVKRRLDVNIAQGESILEH YRKLGLVTDIEG

#### >SEQ ID NO 2156: 114 CJB110 frame: 1

NLLTTGLLGAGKGTQAAKIVEEFGVAHISTGDMFRAAMANQTEMGRLAKSYIDKGELVPD EVTNGIVKERLAEDDIAEKGFLLDGYPRTIEQAHALDATLEELGLRLDGVINIKVDPSCL IERLSGRIINRKTGETFHKVFNPPVDYKEEDYYQREDDKPETVKRRLDVNIAQGEPILEH

#### >SEQ ID NO 2157: 114 COH1 frame: 3

LLIMGLPGAGKGTQAAKIVEEFGVAHISTGDMFRAAMANQTQMGRLAKSYIDKGELVPDE VTNGIVKERLAEDDIAEKGFLLDGYPRTIEQAHALDATLEELGLRLDGVINIKVDPTCLI ERLSGRIINRKTGETFHKVFNPPVDYKEEDYYQREDDKPETVKRRLDVNIAQGEPILEHY RKLGLVTDIEGNQEITEVFADVEKALL

#### >SEQ ID NO 2158: 114 H36B frame: 3

GDMFRAAMANQTEMGRLAKS\(\frac{\frac{1}{2}}\)IDKGELVPDEVTNGIVKERLAEDDIAEKGFLLDGYPRTI EQAHALDATLEELGLRLDGVINIKVDPSCLIERLSGRIINRKTGETFHKVFNPPVDYKEE DYYQREDDKPETVKRRLDVNIAQGESILEHYRKLGLVTDIEGNQEITEVFADVEKAL

#### >SEQ ID NO 2159: 114\_JM9130013 frame: 1

NILIMGLPGAGKGTQAAKIVEEFGVAHISTGDMFRAAMANQTEMGRLAKSYIDKGELVPD EVTNGIVKERLAEDDIAEKGFLLDGYPRTIEQAHALDATLEELGLRLDGVINIKVDPSCL IERLSGRIINRKTGETFHKVFNPPVDYKEEDYYQREDDKPETVKRRLDVNIAQGEPILEH YKKLGLVTDIEGN

#### >SEQ ID NO 2160:114\_M732 frame: 1

LLIMGLPGAGKGTQAAKIVEEFGVAHISTGDMFRAAMANQTQMGRLAKSYIDKGELVPDE VTNGIVKERLAEDDIAEKGFLLDGYPRTIEQAHALDATLEELGLRLDGVINIKVDPTCLI ERLSGRIINRKTGETFHKVFNPPVDYKEEDYYQREDDKPETVKRRLDVNIAQGEPILEHY RKLGLVTDIEGNQEITEVFADVEKALLELK

#### >SEQ ID NO 2161: 114\_M781 frame: 1

NLLITGLPGAGKGTQAAKIVEEFGVAHISTGDMFRAAMANQTQMGRLAKSYIDKGELVPD EVTNGIVKERLAEDDIAEKGFLLDGYPRTIEQAHALDATLEELGLRLDGVINIKVDPTCL IERLSGRIINRKTGETFHKVFNPPVDYKEEDYYQREDDKPETVKRRLDVNIAQ

## Table 21: Comparative Sequences relating t SAG0079 (adenylate kinase)

	LLIMGLPGAGKGTQAAKIVEEFGVAHISTGDMFRAAMANQTEMGRLAKSYIDKGELVPD
SEQ2151	GKGTQAAKIVEEFGVAHISTGDMFRAAMANQTEMGRLAKSYIDKGELVPD
_	LLTTGSPGAGKGTQAAKIVEEFGVAHISTGDMFRAAMANQTEMGRLAKSYIDKGELVPD
- · · · · · · · · · · · · · · · · · · ·	LLIMGLPGAGKGTQAAKIVEEFGVAHISTGDMFRAAMANQTEMGRLAKSYIDKGELVPD
	LLIMGLPGAGKGTQAAKIVEEFGVAHISTGDMFRAAMANQTEMGRLAKSYIDKGELVPD
	LLIMGLPGAGKGTQAAKIVEEFGVAHISTGDMFRAAMANQTEMGRLAKSYIDKGELVPD
	LLTTGLLGAGKGTQAAKIVEEFGVAHISTGDMFRAAMANQTEMGRLAKSYIDKGELVPD
	LLIMGLPGAGKGTQAAKIVEEFGVAHISTGDMFRAAMANQTQMGRLAKSYIDKGELVPD
SEQ2158	GDMFRAAMANQTEMGRLAKSYIDKGELVPD
	LLIMGLPGAGKGTQAAKIVEEFGVAHISTGDMFRAAMANQTEMGRLAKSYIDKGELVPD
• · · · · · · · · · · · · · · · · · · ·	LLIMGLPGAGKGTQAAKIVEEFGVAHISTGDMFRAAMANQTQMGRLAKSYIDKGELVPD
SEQ2161	LLITGLPGAGKGTQAAKIVEEFGVAHISTGDMFRAAMANQTQMGRLAKSYIDKGELVPD
SEQ2150	EVTNGIVKERLAEDDIAEKGFLLDGYPRTIEQAHALDATLEELGLRLDGVINIKVDPSCL
SEQ2151	QVTNGIVKERLAEDDIAEKGFLLDGYPRTIEQAHALDATLEELGLRLDGVINIKVDPSCL
SEQ2152	EVTNGIVKERLAEDDIAEKGFLLDGYPRTIEQAHALDATLEELGLRLDGVINIKVDPSCL
SEQ2153	EVTNGIVKERLAEDDIAEKGFLLDGYPRTIEQAHALDATLEELGLRLDGVINIKVDPSCL
SEQ2154	EVTNGIVKERLAEDDIAEKGFLLDGYPRTIEQAHALDATLEELGLRLDGVINIKVDPSCL.
SEQ2155	EVTNGIVKERLAEDDIAEKGFLLDGYPRTIEQAHALDATLEELGLRLDGVINIKVDPSCL
SEQ2156	EVTNGIVKERLAEDDIAEKGFLLDGYPRTIEQAHALDATLEELGLRLDGVINIKVDPSCL
SEQ2157	EVTNGIVKERLAEDDIAEKGFLLDGYPRTIEQAHALDATLEELGLRLDGVINIKVDPTCL
SEQ2158	EVTNGIVKERLAEDDIAEKGFLLDGYPRTIEQAHALDATLEELGLRLDGVINIKVDPSCL
SEQ2159	EVTNGIVKERLAEDDIAEKGFLLDGYPRTIEQAHALDATLEELGLRLDGVINIKVDPSCL
SEQ2160	EVTNGIVKERLAEDDIAEKGFLLDGYPRTIEQAHALDATLEELGLRLDGVINIKVDPTCL
SEQ2161	EVTNGIVKERLAEDDIAEKGFLLDGYPRTIEQAHALDATLEELGLRLDGVINIKVDPTCL
SEQ2150	IERLSGRIINRKTGETFHKVFNPPVDYKEEDYYQREDDKPETVKRRLDVNIAQGEPILEH
SEQ2151	IERLSGRIINRKTGETFHKVFNPPVDYKEEDYYQREDDKPETVKRRLDVHIAQGEPILEH
SEQ2152	IERLSGRIINRKTGETFHKVFNPPVDYKEEDYYQREDDKPETVKRRLDVNIAQGEPILEH
SEQ2153	IERLSGRIINRKTGETFHKVFNPPVDYKEEDYYQREDDKPETVKRRLDVNIAQGEPILEH
SEQ2154	IERLSGRIINRKTGETFHKVFNPPVDYKEEDYYQREDDKPETVKRRLDVNIAQGESILEH
SEQ2155	IERLSGRIINRKTGETFHKVFNPPVDYKEEDYYQREDDKPETVKRRLDVNIAQGESILEH
SEQ2156	IERLSGRIINRKTGETFHKVFNPPVDYKEEDYYQREDDKPETVKRRLDVNIAQGEPILEH
SEQ2157	IERLSGRIINRKTGETFHKVFNPPVDYKEEDYYQREDDKPETVKRRLDVNIAQGEPILEH
SEQ2158	IERLSGRIINRKTGETFHKVFNPPVDYKEEDYYQREDDKPETVKRRLDVNIAQGESILEH
SEQ2159	IERLSGRIINRKTGETFHKVFNPPVDYKEEDYYQREDDKPETVKRRLDVNIAQGEPILEH
SEQ2160	IERLSGRIINRKTGETFHKVFNPPVDYKEEDYYQREDDKPETVKRRLDVNIAQGEPILEH
SEQ2161	IERLSGRIINRKTGETFHKVFNPPVDYKEEDYYQREDDKPETVKRRLDVNIAQ
SEQ2150	RKLGLVTDIEGNQEITEVFADVEKALLELK
SEQ2151	SKLGLVTDIEGNQEI
SEQ2152	RKLGLVTDIEGNQEITEVFADVEKALLE
SEQ2153	RKLGLVTDIEGNQEITEVFADVEKAL
SEQ2154	RKLGLVTDIEG
SEQ2155	RKLGLVTDIEG
SEQ2156	
SEQ2157	RKLGLVTDIEGNQEITEVFADVEKALL
SEQ2158	RKLGLVTDIEGNQEITEVFADVEKAL
SEQ2159	KKLGLVTDIEGN
SEQ2160	RKLGLVTDIEGNQEITEVFADVEKALLELK
SEQ2161	

# Table 22: Comparative Sequences relating to SAG0093 (D-alanyl-D-alanin carboxypeptidase family protein)

# Table 22: Comparative Sequences relating to SAG0093 (D-alanyl-D-alanine carboxypeptidase family protein)

GGTCAAACATCATTTAACATTAGAAGAATACATAACTTTATTAAAGGAGAATAACCAAAACCCAGCTTTCTTGTACAA

SEQ2201	AGCCTAACAGTCAACAATCATCATCTCAAAAGTTGAGGAATGAGGATATAAAAAAGATA
SEQ2202	AGCCTAACAGTCAACAATCATCACCTCAAAAGTTGAGGAATGAGGATATAAAAAAGATA
SEQ2203	AGCCTAACAGTCAACAATCATCTCCAAAAGTTGAGGAATGAGGATATAAAAAAGATA
SEQ2204	ACAGTCAACAATCATCATCTCAAAAGTTGAGGAATGAGGATATAAAAAAGATA
SEQ2205	AGCCTAACAGTCAACAATCATCTCAAAAGTTGAGGAATGAGGATATAAAAAAGACA
SEQ2206	AGCCTAACAGTCAACAATCATCTCAAAAGTTGAGGAATGAGGATATAAAAAAGATA
SEQ2207	CCTAACAGTCAACAATCATCATCTCAAAAGTTGAGGAATGAGGATATAAAAAAGACA
SEQ2208	AGCCTAACAGTCAACAATCATCTCAAAAGTTGAGGAATGAGGATATAAAAAAGACA
SEQ2209	AGCCTAACAGTCAACAATCATCATCTCAAAAGTTGAGGAATGAGGATATAAAAAAGATA
SEQ2210	AGCCTAACAGTCAACAATCATCTCAAAAGTTGAGGAATGAGGATATAAAAAAGACA
SEQ2211	AGCCTAACAGTCAACAATCATCTCAAAAGTTGAGGAATGAGGATATAAAAAAGACA

# Table 22: Comparative Sequences relating to SAG0093 (D-alanyl-D-alanine carboxypeptidase family protein)

SEQ2201	TCCTCTCAAAAAGAAAT-AAGAAATT-ACAATTACCAGCTGTATCATCAAAAGATTGGA
SEQ2202	TCCTCTCAAAAAGAAT-AAGAAATT-ACGATTACCAGCTGTATCATCAAAAGATTGGA
SEQ2203	TCCTCTCAAAAAAGAAAT-AAGAAATT-ACAATTACCAGCTGTATCATCAAAAGATTGGA
SEQ2204	TCCTCTCAAAAAAGAAAT-AAGAAATT-ACAATTACCAGCTGTATCATCAAAAGATTGGA
SEQ2205	TCCTCTCAAAAAAGAAAT-AAGAAATT-ACGATTACCAGCTGTATCATCAAAAGATTGGA
SEQ2206	TCCTCTCAAAAAAGAAAT-AAGAAATTTACAATTACCAGCTGTATCATCAAAAGATTGGA
SEQ2207	TCCTCTCAAAAAAGAAATTAAGAAATT-ACGATTACCAGCTGTATCATCAAAAGATTGGA
SEQ2208	TCCTCTCAAAAAGAAAT-AAGAAATT-ACGATTACCAGCTGTATCATCAAAAGATTGGA
SEQ2209	TCCTCTCAAAAAGAAAT-AAGAAATT-ACAATTACCAGCTGTATCATCAAAAGATTGGA
SEQ2210	TCCTCTCAAAAAAGAAAT-AAGAAATT-ACGATTACCAGCTGTATCATCAAAAGATTGGA
SEQ2211	TCCTCTCAAAAAAGAAAT-AAGAAATT-ACGATTACCAGCTGTATCATCAAAAGATTGGA
DDZCCTT	TOTOTOTEWARMOREM PROFESSION FOR THE PROFESSION FOR
SEQ2201	ACTTGATTTTGGTCAATCGTGACCATAAACATGAAGAATTAAGTCCAGATGTGGTTCCTG
SEQ2202	ACTTGATTTTGGTCAATCGTGACCATAAACATGAAGAATTAAGTCCAGATGTGGTGCCTG
SEQ2203	ACTTGATTTTGGTCAATCGTGACCATAAACATGAAGAATTAAGTCCAGATGTGGTTCCTG
SEQ2204	ACTTGATTTTGGTCAATCGTGACCATAAACATGAAGAATTAAGTCCAGATGTGGTTCCTG
SEQ2205	ACTIGATITIGGICAATCGIGACCATAAACATGAAGAATTAAGTCCAGATGTGGTGCCTG
SEQ2205	ACTIGATITIGGICAATCGIGACCATAAACATGAAGAATTAAGTCCAGATGIGGTTCCTG
SEQ2207	ACTIGATITIGGICAATCGIGACCATAAACATGAAGAATTAAGICCAGATGIGGITCCIG ACTTGATTTTGGTCAATCGTGACCATAAACATGAAGAATTAAGICCAGATGIGGTGCCTG
<del></del>	
SEQ2208	ACTTGATTTTGGTCAATCGTGACCATAAACATGAAGAATTAAGTCCAGATGTGGTGCCTG
SEQ2209	ACTTGATTTTGGTCAATCGTGACCATAAACATGAAGAATTAAGTCCAGATGTGGTTCCTG
SEQ2210	ACTTGATTTTGGTCAATCGTGACCATAAACATGAAGAATTAAGTCCAGATGTGGTGCCTG
SEQ2211	. ACTTGATTTTGGTCAATCGTGACCATAAACATGAAGAATTAAGTCCAGATGTGGTGCCTG
SEQ2201	TTGAAAATATTTATTTGGATAAACGTATTACGAAGCAAGC
SEQ2201	TTGAAAATATTTATTTGGATAAACGTATTACGAAGCAAGC
SEQ2202 SEQ2203	TTGAAAATATTTATTTGGATAAACGTATTACGAAGCAAGC
	TTGAAAATATTTATTTGGATAAACGTATTACGAAGCAAGC
SEQ2204	
SEQ2205 SEQ2206	TTGAAAATATTTATTTGGATAAACGTATTACGAAGCAAGC
SEQ2208	TTGAAAATATTTATTTGGATAAACGTATTACGAAGCAAGC
SEQ2207 SEQ2208	
_	TTGAAAATATTTATTTGGATAAACGTATTACGAAGCAAGC
SEQ2209	TTGAAAATATTTATTTGGATAAACGTATTACGAAGCAAGC
SEQ2210	TTGAAAATATTTATTTGGATAAACGTATTACGAAGCAAGC
SEQ2211	TTGAAAATATTTATTTGGATAAACGTATTACGAAGCAAGC
SEQ2201	CTAGAGCAATTGATTCACGAGAACATTTAATTTCGGGTTATCGTAGTGTTGCCTATCAGG
SEQ2202	CTAGAGCAATTGATTCACGAGAACATTTAATTTCGGGTTATCGTAGTGTTGCCTATCAGG
SEQ2203	CTAGAGCAATTGATTCACGAGAACATTTAATTTCGGGTTATCGTAGTGTTGCCTATCAGG
SEQ2204	CTAGAGCAATTGATTCACGAGAACATTTAATTTCGGGTTATCGTAGTGTTGCCTATCAGG
SEQ2205	CTAGAGCAATTGATTCACGAGAACATTTAATTTCGGGTTATCGTAGTGTTGCCTATCAGG
SEQ2205 SEQ2206	CTAGAGCAATTGATTCACGAGAACATTTAATTTCGGGTTATCGTAGTGTTGCCTATCAGG CTAGAGCAATTGATTCACGAGAACATTTAATTTCGGGTTATCGTAGTGTTGCCTATCAGG
SEQ2200 SEQ2207	CTAGAGCAATTGATTCACGAGAACATTTAATTTCGGGTTATCGTAGTGTTGCCTATCAGG CTAGAGCAATTGATTCACGAGAACATTTAATTTCGGGTTATCGTAGTGTTGCCTATCAGG
SEQ2207 SEQ2208	CTAGAGCAATTGATTCACGAGAACATTTAATTTCGGGTTATCGTAGTGTTGCCTATCAGG
SEQ2208 SEQ2209	CTAGAGCAATTGATTCACGAGAACATTTAATTTCGGGTTATCGTAGTGTTGCCTATCAGG
SEQ2209 SEQ2210	CTAGAGCAATTGATTCACGAGAACATTTAATTTCGGGTTATCGTAGTGTTGCCTATCAGG
SEQ2210 SEQ2211	CTAGAGCAATTGATTCACGAGAACATTTAATTTCGGGTTATCGTAGTGTTGCCTATCAGG CTAGAGCAATTGATTCACGAGAACATTTAATTTCGGGTTATCGTAGTGTTGCCTATCAGG
2505511	CHAGAGCAMITGATICACGAGAACATTAATTICGGGITATCGTAGTGTTGCCTATCAGG
SE02201	AGAAGTTGTTCAATTCTTATGTTACTCAAGAGATGACTAGTAACCCTAATTTGACGAGGG
SEQ2202	AGAAGTTGTTCAATTCTTATGTTACTCAAGAGATGACTAGTAACCCTAATTTGACGAGGG
SEQ2202 SEQ2203	AGAAGTTGTTCAATTCTTATGTTACTCAAGAGATGACTAGTAACCCTAATTTGACGAGGG
SEQ2203 SEQ2204	AGAAGTTGTTCAATTCTTATGTTACTCAAGAGATGACTAGTAACCCTAATTTGACGAGGG AGAAGTTGTTCAATTCTTATGTTACTCAAGAGATGACTAGTAACCCTAATTTGACGAGGG
SEQ2204 SEQ2205	AGAAGTTGTTCAATTCTTATGTTACTCAAGAAATGACTAGTAACCCTAATTTGACGAGGG AGAAGTTGTTCAATTCTTATGTTACTCAAGAAATGACTAGTAACCCTAATTTGACGAAGG
SEQ2206	AGAAGTTGTTCAATTCTTATGTTACTCAAGAGATGACTAGTAACCCTAATTTGACGAGGG AGAAGTTGTTCAATTCTTATGTTACTCAAGAGATGACTAGTAACCCTAATTTGACGAGGG
SEQ2207	AGAAGI I GTTCAATTCTTATGTTACTCAAGAGATGACTAGTAACCCTAATTTGACGAGGG
SEQ2208	AGAAGTTGTTCAATTCTTATGTTACTCAWGAAATGACTAGTAACCCTAATTTGACGAAGG
SEQ2209	AGAAGTTGTTCAATTCTTATGTTACTCAAGAGATGACTAGTAACCCTAATTTGACGAGGG
SEQ2210	AGAAGTTGTTCAATTCTTATGTTACTCAAGAGATGACTAGTAACCCTAATTTGACGAGGG
SEQ2211	AGAAGTTGTTCAATTCTTATGTTACTCAAGAGATGACTAGTAACCCTAATTTGACGAGGG

# Table 22: Comparative Sequences relating to SAG0093 (D-alanyl-D-alanine carb xypeptidase family protein)

SEQ2201	acaagcagaaaagttggtaaaaacttactctcagcctgcaggtgctagtgaacaccaga
SEQ2202	acaagcagaaaagttggtaaaaacttactctcagcctgcaggtgctagtgaacaccaga
SEQ2203	acaagcagaaagttggtaaaaacttactctcagcctgcaggtgctagtgaacaccaga
SEQ2204	acaagcagaaagttggtaaaaacttactctcagcctgcaggtgctagtgaacaccaga
SEQ2205	ACAAGCAGAAAAGTTGGTAAAAACTTACTCTCAGCCTGCAGGTGCTAGTGAACACCAGA
SEQ2206	ACAAGCAGAAAAGTTGGTAAAAACTTACTCTCAGCCTGCAGGTGCTAGTGAACACCAGA
SEQ2207	ACAAGCAGAAAAGTTGGTAAAAACTTACTCTCAGCCTGCAGGTGCTAGTGAACACCAGA
SEQ2208	ACAAGCAGAAAAGTTGGTAAAAACTTACTCTCAGCCTGCAGGTGCTAGTGAACACCAGA
SEQ2209	ACAAGCAGAAAAGTTGGTAAAAACTTACTCTCAGCCTGCAGGTGCTAGTGAACACCAGA
SEQ2210	ACAAGCAGAAAAGTTGGTAAAAACTTACTCTCAGCCTGCAGGTGCTAGTGAACACCAGA
SEQ2211	ACAAGCAGAAAAGTTGGTAAAAACTTACTCTCAGCCTGCAGGTGCTAGTGAACACCAGA
DDZCCTT	
SEQ2201	CTGGATTAGCGATGGATATGAGTACTGTAGATTCTTTGAATGAGAGCGATCCTAGAGTAG
SEQ2202	CTGGATTAGCGATGGATATGAGTACTGTAGATTCTTTGAATGAGAGCGATCCTAGAGTAG
SEQ2203 ·	CTGGATTAGCGATGGATATGAGTACTGTAGATTCTTTGAATGAGAGCGATCCTAGAGTAG
SEQ2204	CTGGATTAGCGATGGATATGAGTACTGTAGATTCTTTGAATGAGAGCGATCCTAGAGTAG
SEQ2205	CTGGATTAGCGATGGATATGAGTACTGTAGATTCTTTGAATGAGAGCGATCCTAGAGTAG
SEQ2205 SEQ2206	CTGGATTAGCGATGGATATGAGTACTGTAGATTCTTTGAATGAGAGCGATCCTAGAGTAG
SEQ2206 SEQ2207	CTGGATTAGCGATGGATATGAGTACTGTAGATTCTTTGAATGAGAGCGATCCTAGAGTAG
-	CTGGATTAGCGATGGATATGAGTACTGTAGATTCTTTGAATGAGAGCGATCCTAGAGTAG
SEQ2208	CTGGATTAGCGATGGATATGAGTACTGTAGATTCTTTGAATGAGGCGATCCTAGAGTAG
SEQ2209	CTGGATTAGCGATGGATATGAGTACTGTAGATTCTTTGAATGAGAGCGATCCTAGAGTAG
SEQ2210	CTGGATTAGCGATGGATATGAGTACTGTAGATTCTTTGAATGAGAGCGATCCTAGAGTAG
SEQ2211	CTGGATTAGCGATGGATATGAGTACTGTAGATTCTTTGAATGAGCGATCCTAGAGTAG
SEQ2201	TCAGTCAGTTGAAAAAGATAGCTCCACAATATGGTTTTGTCTTACGGTTTCCGGATGGTA
-	TCAGTCAGTTGAAAAAGATAGCTCCACAATATGGTTTTGTCTTACGGTTTCCGGATGGTA
SEQ2202	TCAGTCAGTTGAAAAAGATAGCTCCACAATATGGTTTTGTCTTACGGTTTCCGGATGGTA
SEQ2203	TCAGTCAGTTGAAAAAGATAGCTCCACAATATGGTTTTGTCTTACGGTTTCCGGATGGTA TCAGTCAGTTGAAAAAGATAGCTCCACAATATGGTTTTGTCTTACGGTTTCCGGATGGTA
SEQ2204	TCAGTCAGTTGAAAAAGATAGCTCCACAATATGGTTTTGTCTTACGGTTTCCGGATGGTA TCAGTCAGTTGAAAAAGATAGCTCCACAATATGGTTTTGTCTTACGGTTTCCGGATGGTA
SEQ2205	TCAGTCAGTTGAAAAAGATAGCTCCACAATATGGTTTTGTCTTACGGTTTCCGGATGGTA TCAGTCAGTTGAAAAAGATAGCTCCACAATATGGTTTTGTCTTACGGTTTCCGGATGGTA
SEQ2206	TCAGTCAGTTGAAAAAGATAGCTCCACAATATGGTTTTGTCTTACGGTTTCCGGATGGTA TCAGTCAGTTGAAAAAGATAGCTCCACAATATGGTTTTGTCTTACGGTTTCCGGATGGTA
SEQ2207	
SEQ2208	TCAGTCAGTTGAAAAAGATAGCTCCACAATATGGTTTTGTCTTACGGTTTCCGGATGGTA
SEQ2209	TCAGTCAGTTGAAAAAGATAGCTCCACAATATGGTTTTGTCTTACGGTTTCCGGATGGTA
SEQ2210	TCAGTCAGTTGAAAAAGATAGCTCCACAATATGGTTTTGTCTTACGGTTTCCGGATGGTA
SEQ2211	TCAGTCAGTTGAAAAAGATAGCTCCACAATATGGTTTTGTCTTACGGTTTCCGGATGGTA
SEQ2201	AAACAGCAGAAACAGGGGTAGGTTATGAAGATTGGCATTACCGCTATGTTGGGGTAGAGT
SEQ2201	AAACAGCAGAAACAGGGGTAGGTTATGAAGATTGGCATTACCGCTATGTTGGGGTAGAGT
SEQ2202	AAACAGCAGAAACAGGGGTAGGTTATGAAGATTGGCATTACCGCTATGTTGGGGTAGAGT
SEQ2204	AAACAGCAGAAACAGGGGTAGGTTATGAAGATTGGCATTACCGCTATGTTGGGGTAGAGT
SEQ2205	AAACAGCAGAAACAGGGGTAGGTTATGAAGATTGGCATTACCGCTATGTTGGGGTAGAGT
SEQ2205 SEQ2206	AAACAGCAGAAACAGGGGTAGGTTATGAAGATTGGCATTACCGCTATGTTGGGGTAGAGT
SEQ2207	AAACAGCAGAAACAGGGGTAGGTTATGAAGATTGGCATTACCGCTATGTTGGGGTAGAGT
SEQ2207 SEQ2208	AAACAGCAGAAACAGGGGTAGGTTATGAAGATTGGCATTACCGCTATGTTGGGGTAGAGT
SEQ2208 SEQ2209	AAACAGCAGAAACAGGGGTAGGTTATGAAGATTGGCATTACCGCTATGTTGGGGTAGAGT AAACAGCAGAAACAGGGGTAGGTTATGAAGATTGGCATTACCGCTATGTTGGGGTAGAGT
_	AAACAGCAGAAACAGGGGTAGGTTATGAAGATTGGCATTACCGCTATGTTGGGGTAGAGT
SEQ2210 SEQ2211	AAACAGCAGAAACAGGGGTAGGTTATGAAGATTGGCATTACCGCTATGTTGGGGTAGAGT AAACAGCAGAAACAGGGGTAGGTTATGAAGATTGGCATTACCGCTATGTTGGGGTAGAGT
SEGSSII	ANACAGEAGANACAGGGTAGGTATT GAAGATTAGGGTATGT TOCCOTAGAT
SEQ2201	CTGCAAAATATATGGCCAAACATCATTTAACATTAGAAGAATACATAACTTTATTAAAGG
SEQ2202	CTGCAAAATATATGGCCGAACATCGTTTAACATTAGAAGAATACATAACTTTATTAAAGG
SEQ2203	CTGCAAAATATATGGCCAAACATCATTTAACATTAGAAGAATACATAACTTTATTAAAGG
SEQ2204	CTGCAAAATATATGGCCAAACATCATTTAACATTAGAAGAATACATTAACTTTATTAAAGG
SEQ2204 SEQ2205	CTGCAAAATATATGGCCAAACATCATTAACATTAGAAGAATACATAACTTTATTAAAGG
SEQ2205 SEQ2206	CTGCAAAATATATGGCCAAACATCATTTAACATTAGAAGAATACATTAACTTTATTAAAGG
SEQ2206 SEQ2207	CTGCAAAATATATGGTCAAACATCATTTAACATTAGAAGAATACATTATAACATTATAAAGG
-	CTGCAAAATATATGGTCAAACATCATTTAACATTAGAAGAATACATTAACTTTATTAAAAGG CTGCAAAATATATGGCCAAACATCATTTAACATTAGAAGAATACATTAACTTTATTAAAAGG
SEQ2208	CTGCAAAATATATGGCCAAACATCATTTAACATTAGAAGAATACATAACTTTATTAAAAGG CTGCAAAATATATGGCCAAACATCATTTAACATTAGAAGAATACATAACTTTATTAAAAGG
SEQ2209	CTGCAAAATATATGGCCAAACATCATTTAACATTAGAAGAATACATTAACTTTATTAAAGG CTGCAAAATATATGGTCAAACATCATTTAACATTAGAAGAATACATAACTTTATTAAAGG
SEQ2210	CTGCAAAATATATGGTCAAACATCATTTAACATTAGAAGAATACATTAACTTTATTAAAGG CTGCAAAATATATGGTCAAACATCATTTAACATTAGAAGAATACATTAACTTTATTAAAAGG
SEQ2211	CIGCAAAMIAIAIGGICAAACAICAITIAACAITAGAAGAATACAIAACTITAITAAAGG



# Table 22: Comparative Sequences relating to SAG0093 (D-alanyl-D-alanine carb xypeptidase family protein)

SEQ2201	AGAATAACCAA
SEQ2202	AGAATAACCAA
SEQ2203	AGAATAACCAA
SEQ2204	AGAATAACCAAAACCCAGCTTTCTTGTACAA
SEQ2205	AGAATAACCAA
SEQ2206	AGAATAACCAA
SEQ2207	AGAATAACCAAAACCCAGCTTTCTTGTACAA
SEQ2208	AGAATAACCAA
SEQ2209	AGAATAACCAA
SEQ2210	AGAATAACCAAAACCCAGCTTTCTT
SEQ2211	AGAATAACCAATABCMARATVSTNCSRATNGTSAGDAANYDAANNCARBXYTDASAMYRT

#### >SEQ ID NO 2250: 18 090 frame: 1

KPNSQQSSSQKLRNEDIKKISSQKRNKKLQLPAVSSKDWNLILVNRDHKHEELSPDVVPV ENIYLDKRITKQATQFLEAARAIDSREHLISGYRSVAYQEKLFNSYVTQEMTSNPNLTRG QAEKLVKTYSQPAGASEHQTGLAMDMSTVDSLNESDPRVVSQLKKIAPQYGFVLRFPDGK TAETGVGYEDWHYRYVGVESAKYMAKHHLTLEEYITLLKENNQ

#### >SEQ ID NO 2251: 18 1169NT frame: 1

KPNSQQSSPQKLRNEDIKKISSQKRNKKLRLPAVSSKDWNLILVNRDHKHEELSPDVVPV ENIYLDKRITKQATQFLEAARAIDSREHLISGYRSVAYQEKLFNSYVTQEMTSNPNLTRG QAEKLVKTYSQPAGASEHQTGLAMDMSTVDSLNESDPRVVSQLKKIAPQYGFVLRFPDGK TAETGVGYEDWHYRYVGVESAKYMAEHRLTLEEYITLLKENNQ

#### >SEQ ID NO 2252: 18\_18RS21 frame: 1

KPNSQQSSSQKLRNEDIKKISSQKRNKKLQLPAVSSKDWNLILVNRDHKHEELSPDVVPV ENIYLDKRITKQATQFLEAARAIDSREHLISGYRSVAYQEKLFNSYVTQEMTSNPNLTRG QAEKLVKTYSQPAGASEHQTGLAMDMSTVDSLNESDPRVVSQLKKIAPQYGFVLRFPDGK TAETGVGYEDWHYRYVGVESAKYMAKHHLTLEEYITLLKENNQ

#### >SEQ ID NO 2253: 18\_2603 frame: 3

SQQSSSQKLRNEDIKKISSQKRNKKLQLPAVSSKDWNLILVNRDHKHEELSPDVVPVENI YLDKRITKQATQFLEAARAIDSREHLISGYRSVAYQEKLFNSYVTQEMTSNPNLTRGQAE KLVKTYSQPAGASEHQTGLAMDMSTVDSLNESDPRVVSQLKKIAPQYGFVLRFPDGKTAE TGVGYEDWHYRYVGVESAKYMAKHHLTLEEYITLLKENNQNPAFLY

#### >SEQ ID NO 2254: 18\_A909 frame: 1

KPNSQQSSSQKLRNEDIKKTSSQKRNKKLRLPAVSSKDWNLILVNRDHKHEELSPDVVPV ENIYLDKRITKQATQFLEAARAIDSREHLISGYRSVAYQEKLFNSYVTQEMTSNPNLTKE QAEKLVKTYSQPAGASEHQTGLAMDMSTVDSLNESDPRVVSQLKKIAPQYGFVLRFPDGK TAETGVGYEDWHYRYVGVESAKYMAKHHLTLEEYITLLKENNQ

#### >SEQ ID NO 2255:18\_CJB110 frame: 1

KPNSQQSSSQKLRNEDIKKISSQKRNKKFTITSCIIKRLELDFGQS

#### >SEQ ID NO 2256:18\_COH1 frame: 1

PNSQQSSSQKLRNEDIKKTSSQKRN

#### >SEQ ID NO 2257: 18 H36B frame: 1

KPNSQQSSSQKLRNEDIKKTSSQKRNKKLRLPAVSSKDWNLILVNRDHKHEELSPDVVPV ENIYLDKRITKQATQFLEAARAIDSREHLISGYRSVAYQEKLFNSYVTXEMTSNPNLTKE QAEKLVKTYSQPAGASEHQTGLAMDMSTVDSLNESDPRVVSQLKKIAPQYGFVLRFPDGK TAETGVGYEDWHYRYVGVESAKYMAKHHLTLEEYITLLKENNQ

#### >SEQ ID NO 2258: 18\_JM9130013 frame: 1

KPNSQQSSSQKLRNEDIKKISSQKRNKKLQLPAVSSKDWNLILVNRDHKHEELSPDVVPV ENIYLDKRITKQATQFLEAARAIDSREHLISGYRSVAYQEKLFNSYVTQEMTSNPNLTRG QAEKLVKTYSQPAGASEHQTGLAMDMSTVDSLNESDPRVVSQLKKIAPQYGFVLRFPDGK TAETGVGYEDWHYRYVGVESAKYMAKHHLTLEEYITLLKENNQ

#### >SEQ ID NO 2259:18\_M732 frame: 3

PNSQQSSSQKLRNEDIKKTSSQKRNKKLRLPAVSSKDWNLILVNRDHKHEELSPDVVPVE NIYLDKRITKQATQFLEAARAIDSREHLISGYRSVAYQEKLFNSYVTQEMTSNPNLTRGQ AEKLVKTYSQPAGASEHQTGLAMDMSTVDSLNESDPRVVSQLKKIAPQYGFVLRFPDGKT AETGVGYEDWHYRYVGVESAKYMVKHHLTLEEYITLLKENNQNPAF



>SEQ ID NO 2260: 18 M781 frame: 1
KPNSQQSSSQKLRNEDIKKTSSQKRNKKLRLPAVSSKDWNLILVNRDHKHEELSPDVVPV
ENIYLDKRITKQATQFLEAARAIDSREHLISGYRSVAYQEKLFNSYVTQEMTSNPNLTRG
QAEKLVKTYSQPAGASEHQTGLAMDMSTVDSLNESDPRVVSQLKKIAPQYGFVLRFPDGK
TAETGVGYEDWHYRYVGVESAKYMVKHHLTLEEYITLLKENNQ

SEQ2250	PNSQQSSSQKLRNEDIKKISSQKRNKKLQLPAVSSKDWNLILVNRDHKHEELSPDVVPV
SEQ2251	PNSQQSSPQKLRNEDIKKISSQKRNKKLRLPAVSSKDWNLILVNRDHKHEELSPDVVPV
SEQ2252	PNSQQSSSQKLRNEDIKKISSQKRNKKLQLPAVSSKDWNLILVNRDHKHEELSPDVVPV
SEQ2253	SQQSSSQKLRNEDIKKISSQKRNKKLQLPAVSSKDWNLILVNRDHKHEELSPDVVPV
_	
SEQ2254	PNSQQSSSQKLRNEDIKKTSSQKRNKKLRLPAVSSKDWNLILVNRDHKHEELSPDVVPV
SEQ2255	PNSQQSSSQKLRNEDIKKISSQKRNKKFTITSCIIKRLELDFGQS
SEQ2256	PNSQQSSSQKLRNEDIKKTSSQKRN
SEQ2257	PNSQQSSSQKLRNEDIKKTSSQKRNKKLRLPAVSSKDWNLILVNRDHKHEELSPDVVPV
SEQ2258	PNSQQSSSQKLRNEDIKKISSQKRNKKLQLPAVSSKDWNLILVNRDHKHEELSPDVVPV
SEQ2259	PNSQQSSSQKLRNEDIKKTSSQKRNKKLRLPAVSSKDWNLILVNRDHKHEELSPDVVPV
SEQ2260	PNSQQSSSQKLRNEDIKKTSSQKRNKKLRLPAVSSKDWNLILVNRDHKHEELSPDVVPV
SEQUEDO	EUSÄÄSSSÄYTKUESTIVII SSÄYKVUKUTKTENASSYDAUTTTAUKDUKUESTSEDAALA
77000E0	NAME AND THE OWNERS AND ADDRESS AND ADDRES
SEQ2250	NIYLDKRITKQATQFLEAARAIDSREHLISGYRSVAYQEKLFNSYVTQEMTSNPNLTRG
SEQ2251	NIYLDKRITKQATQFLEAARAIDSREHLISGYRSVAYQEKLFNSYVTQEMTSNPNLTRG
SEQ2252	NIYLDKRITKQATQFLEAARAIDSREHLISGYRSVAYQEKLFNSYVTQEMTSNPNLTRG
SEQ2253	NIYLDKRITKQATQFLEAARAIDSREHLISGYRSVAYQEKLFNSYVTQEMTSNPNLTRG
SEQ2254	NIYLDKRITKQATQFLEAARAIDSREHLISGYRSVAYQEKLFNSYVTQEMTSNPNLTKE
SEQ2255	
SEQ2256	
SEQ2257	NIYLDKRITKQATQFLEAARAIDSREHLISGYRSVAYQEKLFNSYVTXEMTSNPNLTKE
SEQ2258	NIYLDKRITKQATQFLEAARAIDSREHLISGYRSVAYQEKLFNSYVTQEMTSNPNLTRG
SEQ2259	NIYLDKRITKQATQFLEAARAIDSREHLISGYRSVAYQEKLFNSYVTQEMTSNPNLTRG
SEQ2260	NIYLDKRITKQATQFLEAARAIDSREHLISGYRSVAYQEKLFNSYVTQEMTSNPNLTRG
SEQ2250	AEKLVKTYSQPAGASEHQTGLAMDMSTVDSLNESDPRVVSQLKKIAPQYGFVLRFPDGK
SEQ2251	AEKLVKTYSQPAGASEHQTGLAMDMSTVDSLNESDPRVVSQLKKIAPQYGFVLRFPDGK
SEQ2252	AEKLVKTYSQPAGASEHQTGLAMDMSTVDSLNESDPRVVSQLKKIAPQYGFVLRFPDGK
SEQ2253	<b>AEKLVKTYSQPAGASEHQTGLAMDMSTVDSLNESDPRVVSQLKKIAPQYGFVLRFPDGK</b>
SEQ2254	AEKLVKTYSQPAGASEHQTGLAMDMSTVDSLNESDPRVVSQLKKIAPQYGFVLRFPDGK
SEQ2255	
SEQ2256	
SEQ2257	AEKLVKTYSQPAGASEHQTGLAMDMSTVDSLNESDPRVVSQLKKIAPQYGFVLRFPDGK
SEQ2258	AEKLVKTYSQPAGASEHQTGLAMDMSTVDSLNESDPRVVSQLKKIAPQYGFVLRFPDGK
SEQ2258 SEQ2259	AEKLVKTYSQPAGASEHQTGLAMDMSTVDSLNESDPRVVSQLKKIAPQYGFVLRFPDGK AEKLVKTYSQPAGASEHQTGLAMDMSTVDSLNESDPRVVSQLKKIAPQYGFVLRFPDGK
SEQ2258	AEKLVKTYSQPAGASEHQTGLAMDMSTVDSLNESDPRVVSQLKKIAPQYGFVLRFPDGK
SEQ2258 SEQ2259 SEQ2260	AEKLVKTYSQPAGASEHQTGLAMDMSTVDSLNESDPRVVSQLKKIAPQYGFVLRFPDGK AEKLVKTYSQPAGASEHQTGLAMDMSTVDSLNESDPRVVSQLKKIAPQYGFVLRFPDGK AEKLVKTYSQPAGASEHQTGLAMDMSTVDSLNESDPRVVSQLKKIAPQYGFVLRFPDGK
SEQ2258 SEQ2259 SEQ2260 SEQ2250	AEKLVKTYSQPAGASEHQTGLAMDMSTVDSLNESDPRVVSQLKKIAPQYGFVLRFPDGK AEKLVKTYSQPAGASEHQTGLAMDMSTVDSLNESDPRVVSQLKKIAPQYGFVLRFPDGK AEKLVKTYSQPAGASEHQTGLAMDMSTVDSLNESDPRVVSQLKKIAPQYGFVLRFPDGK AETGVGYEDWHYRYVGVESAKYMAKHHLTLEEYITLLKENNQ
SEQ2258 SEQ2259 SEQ2260	AEKLVKTYSQPAGASEHQTGLAMDMSTVDSLNESDPRVVSQLKKIAPQYGFVLRFPDGK AEKLVKTYSQPAGASEHQTGLAMDMSTVDSLNESDPRVVSQLKKIAPQYGFVLRFPDGK AEKLVKTYSQPAGASEHQTGLAMDMSTVDSLNESDPRVVSQLKKIAPQYGFVLRFPDGK
SEQ2258 SEQ2259 SEQ2260 SEQ2250	AEKLVKTYSQPAGASEHQTGLAMDMSTVDSLNESDPRVVSQLKKIAPQYGFVLRFPDGK AEKLVKTYSQPAGASEHQTGLAMDMSTVDSLNESDPRVVSQLKKIAPQYGFVLRFPDGK AEKLVKTYSQPAGASEHQTGLAMDMSTVDSLNESDPRVVSQLKKIAPQYGFVLRFPDGK AETGVGYEDWHYRYVGVESAKYMAKHHLTLEEYITLLKENNQ
SEQ2258 SEQ2259 SEQ2260 SEQ2250 SEQ2251	AEKLVKTYSQPAGASEHQTGLAMDMSTVDSLNESDPRVVSQLKKIAPQYGFVLRFPDGK AEKLVKTYSQPAGASEHQTGLAMDMSTVDSLNESDPRVVSQLKKIAPQYGFVLRFPDGK AEKLVKTYSQPAGASEHQTGLAMDMSTVDSLNESDPRVVSQLKKIAPQYGFVLRFPDGK AETGVGYEDWHYRYVGVESAKYMAKHHLTLEEYITLLKENNQ
SEQ2258 SEQ2259 SEQ2260 SEQ2250 SEQ2251 SEQ2252 SEQ2252	AEKLVKTYSQPAGASEHQTGLAMDMSTVDSLNESDPRVVSQLKKIAPQYGFVLRFPDGK AEKLVKTYSQPAGASEHQTGLAMDMSTVDSLNESDPRVVSQLKKIAPQYGFVLRFPDGK AEKLVKTYSQPAGASEHQTGLAMDMSTVDSLNESDPRVVSQLKKIAPQYGFVLRFPDGK AETGVGYEDWHYRYVGVESAKYMAKHHLTLEEYITLLKENNQ
SEQ2258 SEQ2259 SEQ2260 SEQ2250 SEQ2251 SEQ2252 SEQ2253 SEQ2253	AEKLVKTYSQPAGASEHQTGLAMDMSTVDSLNESDPRVVSQLKKIAPQYGFVLRFPDGK AEKLVKTYSQPAGASEHQTGLAMDMSTVDSLNESDPRVVSQLKKIAPQYGFVLRFPDGK AEKLVKTYSQPAGASEHQTGLAMDMSTVDSLNESDPRVVSQLKKIAPQYGFVLRFPDGK AETGVGYEDWHYRYVGVESAKYMAKHHLTLEEYITLLKENNQ
SEQ2258 SEQ2259 SEQ2260 SEQ2251 SEQ2251 SEQ2252 SEQ2253 SEQ2254 SEQ2255	AEKLVKTYSQPAGASEHQTGLAMDMSTVDSLNESDPRVVSQLKKIAPQYGFVLRFPDGK AEKLVKTYSQPAGASEHQTGLAMDMSTVDSLNESDPRVVSQLKKIAPQYGFVLRFPDGK AEKLVKTYSQPAGASEHQTGLAMDMSTVDSLNESDPRVVSQLKKIAPQYGFVLRFPDGK AETGVGYEDWHYRYVGVESAKYMAKHHLTLEEYITLLKENNQ
SEQ2258 SEQ2259 SEQ2260 SEQ2250 SEQ2251 SEQ2252 SEQ2253 SEQ2253 SEQ2254 SEQ2255 SEQ2255	AEKLVKTYSQPAGASEHQTGLAMDMSTVDSLNESDPRVVSQLKKIAPQYGFVLRFPDGK AEKLVKTYSQPAGASEHQTGLAMDMSTVDSLNESDPRVVSQLKKIAPQYGFVLRFPDGK AEKLVKTYSQPAGASEHQTGLAMDMSTVDSLNESDPRVVSQLKKIAPQYGFVLRFPDGK AETGVGYEDWHYRYVGVESAKYMAKHHLTLEEYITLLKENNQ
SEQ2258 SEQ2259 SEQ2260 SEQ2251 SEQ2251 SEQ2252 SEQ2253 SEQ2254 SEQ2255 SEQ2255 SEQ2255	AEKLVKTYSQPAGASEHQTGLAMDMSTVDSLNESDPRVVSQLKKIAPQYGFVLRFPDGK AEKLVKTYSQPAGASEHQTGLAMDMSTVDSLNESDPRVVSQLKKIAPQYGFVLRFPDGK AEKLVKTYSQPAGASEHQTGLAMDMSTVDSLNESDPRVVSQLKKIAPQYGFVLRFPDGK AETGVGYEDWHYRYVGVESAKYMAKHHLTLEEYITLLKENNQ
SEQ2258 SEQ2259 SEQ2260 SEQ2250 SEQ2251 SEQ2252 SEQ2253 SEQ2254 SEQ2254 SEQ2255 SEQ2256 SEQ2256 SEQ2256	AEKLVKTYSQPAGASEHQTGLAMDMSTVDSLNESDPRVVSQLKKIAPQYGFVLRFPDGK AEKLVKTYSQPAGASEHQTGLAMDMSTVDSLNESDPRVVSQLKKIAPQYGFVLRFPDGK AEKLVKTYSQPAGASEHQTGLAMDMSTVDSLNESDPRVVSQLKKIAPQYGFVLRFPDGK AETGVGYEDWHYRYVGVESAKYMAKHHLTLEEYITLLKENNQ
SEQ2258 SEQ2259 SEQ2250 SEQ2251 SEQ2251 SEQ2252 SEQ2253 SEQ2254 SEQ2254 SEQ2255 SEQ2256 SEQ2256 SEQ2256 SEQ2257	AEKLVKTYSQPAGASEHQTGLAMDMSTVDSLNESDPRVVSQLKKIAPQYGFVLRFPDGK AEKLVKTYSQPAGASEHQTGLAMDMSTVDSLNESDPRVVSQLKKIAPQYGFVLRFPDGK AEKLVKTYSQPAGASEHQTGLAMDMSTVDSLNESDPRVVSQLKKIAPQYGFVLRFPDGK AETGVGYEDWHYRYVGVESAKYMAKHHLTLEEYITLLKENNQ
SEQ2258 SEQ2259 SEQ2260 SEQ2250 SEQ2251 SEQ2252 SEQ2253 SEQ2254 SEQ2254 SEQ2255 SEQ2256 SEQ2256 SEQ2256	AEKLVKTYSQPAGASEHQTGLAMDMSTVDSLNESDPRVVSQLKKIAPQYGFVLRFPDGK AEKLVKTYSQPAGASEHQTGLAMDMSTVDSLNESDPRVVSQLKKIAPQYGFVLRFPDGK AEKLVKTYSQPAGASEHQTGLAMDMSTVDSLNESDPRVVSQLKKIAPQYGFVLRFPDGK AETGVGYEDWHYRYVGVESAKYMAKHHLTLEEYITLLKENNQ
SEQ2258 SEQ2259 SEQ2250 SEQ2251 SEQ2252 SEQ2253 SEQ2254 SEQ2255 SEQ2256 SEQ2256 SEQ2257 SEQ2258 SEQ2258 SEQ2259 SEQ2260	AEKLVKTYSQPAGASEHQTGLAMDMSTVDSLNESDPRVVSQLKKIAPQYGFVLRFPDGK AEKLVKTYSQPAGASEHQTGLAMDMSTVDSLNESDPRVVSQLKKIAPQYGFVLRFPDGK AEKLVKTYSQPAGASEHQTGLAMDMSTVDSLNESDPRVVSQLKKIAPQYGFVLRFPDGK AETGVGYEDWHYRYVGVESAKYMAKHHLTLEEYITLLKENNQ
SEQ2258 SEQ2259 SEQ2250 SEQ2251 SEQ2251 SEQ2252 SEQ2253 SEQ2254 SEQ2255 SEQ2256 SEQ2257 SEQ2257 SEQ2258 SEQ2259 SEQ2250 SEQ2250	AEKLVKTYSQPAGASEHQTGLAMDMSTVDSLNESDPRVVSQLKKIAPQYGFVLRFPDGK AEKLVKTYSQPAGASEHQTGLAMDMSTVDSLNESDPRVVSQLKKIAPQYGFVLRFPDGK AEKLVKTYSQPAGASEHQTGLAMDMSTVDSLNESDPRVVSQLKKIAPQYGFVLRFPDGK AETGVGYEDWHYRYVGVESAKYMAKHHLTLEEYITLLKENNQ
SEQ2258 SEQ2259 SEQ2250 SEQ2251 SEQ2252 SEQ2253 SEQ2254 SEQ2255 SEQ2256 SEQ2256 SEQ2257 SEQ2258 SEQ2258 SEQ2259 SEQ2260	AEKLVKTYSQPAGASEHQTGLAMDMSTVDSLNESDPRVVSQLKKIAPQYGFVLRFPDGK AEKLVKTYSQPAGASEHQTGLAMDMSTVDSLNESDPRVVSQLKKIAPQYGFVLRFPDGK AEKLVKTYSQPAGASEHQTGLAMDMSTVDSLNESDPRVVSQLKKIAPQYGFVLRFPDGK AETGVGYEDWHYRYVGVESAKYMAKHHLTLEEYITLLKENNQ
SEQ2258 SEQ2259 SEQ2250 SEQ2251 SEQ2251 SEQ2252 SEQ2253 SEQ2254 SEQ2255 SEQ2256 SEQ2257 SEQ2257 SEQ2258 SEQ2259 SEQ2250 SEQ2250	AEKLVKTYSQPAGASEHQTGLAMDMSTVDSLNESDPRVVSQLKKIAPQYGFVLRFPDGK AEKLVKTYSQPAGASEHQTGLAMDMSTVDSLNESDPRVVSQLKKIAPQYGFVLRFPDGK AEKLVKTYSQPAGASEHQTGLAMDMSTVDSLNESDPRVVSQLKKIAPQYGFVLRFPDGK AETGVGYEDWHYRYVGVESAKYMAKHHLTLEEYITLLKENNQ
SEQ2258 SEQ2259 SEQ2250 SEQ2251 SEQ2251 SEQ2253 SEQ2254 SEQ2255 SEQ2256 SEQ2257 SEQ2256 SEQ2257 SEQ2259 SEQ2250 SEQ2250 SEQ2250	AEKLVKTYSQPAGASEHQTGLAMDMSTVDSLNESDPRVVSQLKKIAPQYGFVLRFPDGK AEKLVKTYSQPAGASEHQTGLAMDMSTVDSLNESDPRVVSQLKKIAPQYGFVLRFPDGK AEKLVKTYSQPAGASEHQTGLAMDMSTVDSLNESDPRVVSQLKKIAPQYGFVLRFPDGK AETGVGYEDWHYRYVGVESAKYMAKHHLTLEEYITLLKENNQ
SEQ2258 SEQ2259 SEQ2250 SEQ2251 SEQ2251 SEQ2252 SEQ2253 SEQ2254 SEQ2255 SEQ2256 SEQ2257 SEQ2257 SEQ2258 SEQ2259 SEQ2250 SEQ2250 SEQ2251 SEQ2251 SEQ2251	AEKLVKTYSQPAGASEHQTGLAMDMSTVDSLNESDPRVVSQLKKIAPQYGFVLRFPDGK AEKLVKTYSQPAGASEHQTGLAMDMSTVDSLNESDPRVVSQLKKIAPQYGFVLRFPDGK AEKLVKTYSQPAGASEHQTGLAMDMSTVDSLNESDPRVVSQLKKIAPQYGFVLRFPDGK AETGVGYEDWHYRYVGVESAKYMAKHHLTLEEYITLLKENNQ
SEQ2258 SEQ2259 SEQ2250 SEQ2251 SEQ2251 SEQ2252 SEQ2253 SEQ2254 SEQ2255 SEQ2256 SEQ2257 SEQ2258 SEQ2259 SEQ2259 SEQ2250 SEQ2250 SEQ2251 SEQ2251 SEQ2251 SEQ2252 SEQ2253 SEQ2253	AEKLVKTYSQPAGASEHQTGLAMDMSTVDSLNESDPRVVSQLKKIAPQYGFVLRFPDGK AEKLVKTYSQPAGASEHQTGLAMDMSTVDSLNESDPRVVSQLKKIAPQYGFVLRFPDGK AEKLVKTYSQPAGASEHQTGLAMDMSTVDSLNESDPRVVSQLKKIAPQYGFVLRFPDGK AETGVGYEDWHYRYVGVESAKYMAKHHLTLEEYITLLKENNQ
SEQ2258 SEQ2259 SEQ2250 SEQ2251 SEQ2251 SEQ2252 SEQ2253 SEQ2254 SEQ2255 SEQ2256 SEQ2257 SEQ2258 SEQ2259 SEQ2250 SEQ2250 SEQ2250 SEQ2251 SEQ2251 SEQ2251 SEQ2252 SEQ2253 SEQ2253 SEQ2254 SEQ2255	AEKLVKTYSQPAGASEHQTGLAMDMSTVDSLNESDPRVVSQLKKIAPQYGFVLRFPDGK AEKLVKTYSQPAGASEHQTGLAMDMSTVDSLNESDPRVVSQLKKIAPQYGFVLRFPDGK AEKLVKTYSQPAGASEHQTGLAMDMSTVDSLNESDPRVVSQLKKIAPQYGFVLRFPDGK AETGVGYEDWHYRYVGVESAKYMAKHHLTLEEYITLLKENNQ
SEQ2258 SEQ2259 SEQ2250 SEQ2251 SEQ2251 SEQ2253 SEQ2254 SEQ2255 SEQ2256 SEQ2257 SEQ2258 SEQ2259 SEQ2259 SEQ2250 SEQ2250 SEQ2250 SEQ2251 SEQ2251 SEQ2252 SEQ2252 SEQ2253 SEQ2253 SEQ2254 SEQ2255 SEQ2255	AEKLVKTYSQPAGASEHQTGLAMDMSTVDSLNESDPRVVSQLKKIAPQYGFVLRFPDGK AEKLVKTYSQPAGASEHQTGLAMDMSTVDSLNESDPRVVSQLKKIAPQYGFVLRFPDGK AEKLVKTYSQPAGASEHQTGLAMDMSTVDSLNESDPRVVSQLKKIAPQYGFVLRFPDGK AETGVGYEDWHYRYVGVESAKYMAKHHLTLEEYITLLKENNQ
SEQ2258 SEQ2259 SEQ2250 SEQ2251 SEQ2251 SEQ2253 SEQ2254 SEQ2255 SEQ2256 SEQ2257 SEQ2258 SEQ2259 SEQ2259 SEQ2250 SEQ2251 SEQ2251 SEQ2251 SEQ2252 SEQ2255 SEQ2255 SEQ2255 SEQ2255 SEQ2255 SEQ2255 SEQ2255	AEKLVKTYSQPAGASEHQTGLAMDMSTVDSLNESDPRVVSQLKKIAPQYGFVLRFPDGK AEKLVKTYSQPAGASEHQTGLAMDMSTVDSLNESDPRVVSQLKKIAPQYGFVLRFPDGK AEKLVKTYSQPAGASEHQTGLAMDMSTVDSLNESDPRVVSQLKKIAPQYGFVLRFPDGK AETGVGYEDWHYRYVGVESAKYMAKHHLTLEEYITLLKENNQ
SEQ2258 SEQ2259 SEQ2250 SEQ2251 SEQ2251 SEQ2253 SEQ2254 SEQ2255 SEQ2256 SEQ2257 SEQ2258 SEQ2259 SEQ2250 SEQ2251 SEQ2251 SEQ2251 SEQ2251 SEQ2251 SEQ2251 SEQ2251 SEQ2253 SEQ2253 SEQ2253 SEQ2254 SEQ2255 SEQ2255 SEQ2256 SEQ2256 SEQ2257 SEQ2258	AEKLVKTYSQPAGASEHQTGLAMDMSTVDSLNESDPRVVSQLKKIAPQYGFVLRFPDGK AEKLVKTYSQPAGASEHQTGLAMDMSTVDSLNESDPRVVSQLKKIAPQYGFVLRFPDGK AEKLVKTYSQPAGASEHQTGLAMDMSTVDSLNESDPRVVSQLKKIAPQYGFVLRFPDGK AETGVGYEDWHYRYVGVESAKYMAKHHLTLEEYITLLKENNQ
SEQ2258 SEQ2259 SEQ2250 SEQ2251 SEQ2251 SEQ2253 SEQ2254 SEQ2255 SEQ2256 SEQ2257 SEQ2258 SEQ2259 SEQ2259 SEQ2250 SEQ2251 SEQ2251 SEQ2251 SEQ2252 SEQ2255 SEQ2255 SEQ2255 SEQ2255 SEQ2255 SEQ2255 SEQ2255	AEKLVKTYSQPAGASEHQTGLAMDMSTVDSLNESDPRVVSQLKKIAPQYGFVLRFPDGK AEKLVKTYSQPAGASEHQTGLAMDMSTVDSLNESDPRVVSQLKKIAPQYGFVLRFPDGK AEKLVKTYSQPAGASEHQTGLAMDMSTVDSLNESDPRVVSQLKKIAPQYGFVLRFPDGK AETGVGYEDWHYRYVGVESAKYMAKHHLTLEEYITLLKENNQ

## Table 23: Comparative Sequences relating to SAG0163 (competence pr tein Cg1A)

## Table 23: Comparative Sequences relating t SAG0163 (competence protein Cg1A)

## Table 23: Comparative Sequences relating to SAG0163 (competence protein Cg1A)



# Table 23: C mparative Sequences relating t SAG0163 (competence protein Cg1A)

SEQ2301	GGCAGTAGAAGTAAATGCTCAAGATATT
SEQ2302	~ #
SEQ2303	TTCAATCATTAGCAAAGCAAGTCATTCATCAGGCAGTAGAAGTAAATGCTCAAGATATT
SEQ2304	GATATT
SEQ2305	TTCAATCATTAGCAAAGCAAGTCATTCATCAGGCAGTAGAAGTAAATGCTCAAGATATT
SEQ2306	TTCAATCATTAGCAAAGCAAGTCATTCATCAGGCAGTAGAAGTAAATGCTCAAGATATT
SEQ2307	
SEQ2308	TCATTAGCAAAGCAAGTCATTCATCAGGCAGTAGAAGTAAATGCTCAAGATATT
SEQ2309	TTCAATCATTAGCAAAGCAAGTCATTCATCAGGCAGTAGAAGTAAATGCTCAAGATATT
SEQ2310	
SEQ2311	CAGTAGAAGTAAATGCTCAAGATATT
SEQ2301	ATATCATTCCCAAAGGTGA-TTGTTATGAA-CTCTATATGCGTATT-GATGATGA
SEQ2302	GGTGA-TTGTTATGAA-ACCTCTACTATTGCGTATTTGATGATGA
SEQ2303	ATATCATTCCCAAAGGTGA-TTGTTATGAA-CTCTATATGCGTATT-GATGATGA
SEQ2304	ATATCATTCCCAAAGGTGA-TTGTTATGAA-CTCTATATGCGTATT-GATGATGA
SEQ2305	ATATCATTCCCAAAGGTGA-TTGTTATGAA-CTCTATATGCGTATT-GATGATGA
SEQ2306	ATATCATTCCCAAAGGTGA-TTGTTATGAA-CTCTATATGCGTATT-GATGATGA
SEQ2307	TGCGTATT-GATGATGA
SEQ2308	ATATCATTCCCAAAGGTGA-TTGTTATGAA-CTCTATATGCGTATT-GATGATGA
SEQ2309	ATATCATTCCCAAAGGTGA-TTGTTATGAA-CTCTATATGCGTATT-GATGATGA
SEQ2310	TGACTTGTTATGAAACTCTATATGCGTATTTGATGATGA
SEQ2311	ATATCATTCCCAAAGGTGA-TTGTTATGAA-TTCTATATGCGTATT-GATGATGA
SEQ2301	AA-GGCGGTTTATTGATGTTTTTGAGTTTAATAGGATGGCTAGTCTTATTAGTCACTTTA
SEQ2302	AA-GGCGGTTTATTGATGTTTTTGAGTTTAATAGGATGGCTAGTCTTATTAGTCACTTTA
SEQ2303	AA-GGCGGTTTATTGATGTTTTTGAGTTTAATAGGATGGCTAGTCTTATTAGTCACTTTA
SEQ2304	AA-GGCGGTTTATTGATGTTTTTGAGTTTAATAGGATGGCTAGTCTTATTAGTCACTTTA
SEQ2305	AA-GGCGGTTTATTGATGTTTTTGAGTTTAATAGGATGGCTAGTCTTATTAGTCACTTTA
SEQ2306	AA-GGCGGTTTATTGATGTTTTTGAGTTTAATAGGATGGCTAGTCTTATTAGTCACTTTA
SEQ2307	AA-GGCGGTTTATTGATGTTTTTGAGTTTAATAGGATGGCTAGTCTTATTAGTCACTTTA
SEQ2308	AA-GGCGGTTTATTGATGTTTTTGAGTTTAATAGGATGGCTAGTCTTATTAGTCACTTTA
SEQ2309	AA-GGCGGTTTATTGATGTTTTTGAGTTTAATAGGATGGCTAGTCTTATTAGTCACTTTA
SEQ2310	AAAGGCGGTTTATTGATGTTTTTGAGTTTAATAGGATGGCTAGTCTTATTAGTCACTTTA
SEQ2311	AA-GGCGGTTTATTGATGTTTTTGAGTTTAATAGGATGGCTAGTCTTATTAGTCACTTTA
SEQ2301	AATTTGTGGCAGGCATGAACGTTGGAGAAAAAAGACGAAGTCAATTAGGTTCTTGTGACT
SEQ2302	AATTTGTGGCAGGCATGAACGTTGGAGAAAAAAGACGAAGTCAATTAGGTTCTTGTGACT
SEQ2303	AATTTGTGGCAGGCATGAACGTTGGAGAAAAAAGACGAAGTCAATTAGGTTCTTGTGACT
SEQ2304	AATTTGTGGCAGGCATGAACGTTGGAGAAAAAAGACGAAGTCAATTAGGTTCTTGTGACT
SEQ2305	AATTTGTGGCAGGCATGAACGTTGGAGAAAAAAGACGAAGTCAATTAGGTTCTTGTGACT
SEQ2306	AATTTGTGGCAGGCATGAACGTTGGAGAAAAAAGACGAAGTCAATTAGGTTCTTGTGACT
SEQ2307	AATTTGTGGCAGGCATGAACGTTGGAGAAAAAAGACGAAGTCAATTAGGTTCTTGTGACT
SEQ2308	AATTTGTGGCAGGCATGAACGTTGGAGAAAAAAGACGAAGTCAATTAGGTTCTTGTGACT
SEQ2309	AATTTGTGGCAGGCATGAACGTTGGAGAAAAAAGACGAAGTCAATTAGGTTCTTGTGACT
SEQ2310	AATTTGTGGCAGGCATGAACGTTGGAGAAAAAAGACGAAGTCAATTAGGTTCTTGTGACT
SEQ2311	AATTTGTGGCAGGCATGAACGTTGGAGAAAAAAGACGAAGTCAATTAGGTTCTTGTGACT

# Tabl 23: Comparative Sequences relating t SAG0163 (competence protein Cg1A)

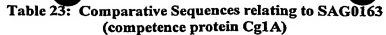
SEQ2301	atgaactgtcagagggaagactggtttcattacgactatcgagtgtgggagattatcgtg
SEQ2302	ATGAACTGTCAGAGGGAAGACTGGTTTCATTACGACTATCGAGTGTGGGAGATTATCGTG
SEQ2303	ATGAACTGTCAGAGGGAAGACTGGTTTCATTACGACTATCGAGTGTGGGAGATTATCGTG
SEQ2304	atgaactgtcagagggaagactggtttcattacgactatcgagtgtgggagattatcgtg
SEQ2305	atgaactgtcagagggaagactggtttcattacgactatcgagtgtgggagattatcgtg
SEQ2306	ATGAACTGTCAGAGGGAAGACTGGTTTCATTACGACTATCGAGTGTGGGAGATTATCGTG
SEQ2307	ATGAACTGTCAGAGGGAAGACTGGTTTCATTACGACTATCAAGTGTGGGAGATTATCGTG
SEQ2308	atgaactgtcagagggaagactggtttcattacgactatcgagtgtgggagattatcgtg
SEQ2309	atgaactgtcagagggaagactggtttcattacgactatcgagtgtgggagattatcgtg
SEQ2310	ATGAACTGTCAGAGGGAAGACTGGTTTCATTACGACTATCAAGTGTGGGGAGATTATCGTG
SEQ2311	ATGAACTGTCAGAGGGAAGACTGGTTTCATTACGACTATCAAGTGTGGGAGATTATCGTG
SEQ2301	GTCAAGAATCTTTAGTTATTCGTATTTTGTATTCAGGTCATCAGGACTTAAAATATTGGT
SEQ2302	GTCAAGAATCTTTAGTTATTCGTATTTTGTATTCAGGTCATCAGGACTTAAAATATTGGT
SEQ2303	GTCAAGAATCTTTAGTTATTCGTATTTTGTATTCAGGTCATCAGGACTTAAAATATTGGT
SEQ2304	GTCAAGAATCTTTAGTTATTCGTATTTTGTATTCAGGTCATCAGGACTTAAAATATTGGT
SEQ2305	GTCAAGAATCTTTAGTTATTCGTATTTTGTATTCAGGTCATCAGGACTTAAAATATTGGT
SEQ2306	GTCAAGAATCTTTAGTTATTCGTATTTTGTATTCAGGTCATCAGGACTTAAAATATTGGT
SEQ2307	GTCAAGAATCTTTAGTTATTCGTACTTTGTATTCAGGTCATCAGGACTTAAAATATTGGT
SEQ2308	GTCAAGAATCTTTAGTTATTCGTATTTTGTATTCAGGTCATCAGGACTTAAAATATTGGT
SEQ2309	GTCAAGAATCTTTAGTTATTCGTATTTTGTATTCAGGTCATCAGGACTTAAAATATTGGT
ŞEQ2310	GTCAAGAATCTTTAGTTATTCGTACTTTGTATTCAGGTCATCAGGACTTAAAATATTGGT
SEQ2311	GTCAAGAATCTTTAGTTATTCGTACTTTGTATTCAGGTCATCAGGACTTAAAATATTGGT
SEQ2301	TTGATAATATAAAGCAAATGAAGGAAGTACTGGGTACAAGAGGGCTATATCTTTTTCCG
SEQ2302	TTGATAATATAAAGCAAATGAAGGAAGTACTGGGTACAAGAGGGCTATATCTTTTTTCCG
SEQ2303	TTGATAATATAAAGCAAATGAAGGAAGTACTGGGTATAAGAGGGCTATATCTTTTTTCCG
SEQ2304	TTGATAATAAAGCAAATGAAGGAAGTACTGGGTATAAGAGGGCTATATCTTTTTTCCG
SEQ2305	TTGATAATATAAAGCAAATGAAGGAAGTACTGGGTATAAGAGGGCTATATCTTTTTTCCG
SEQ2306	TTGATAATATAAAGCAAATGAAGGAAGTACTGGGTACAAGAGGGCTATATCTTTTTTCCG
SEQ2307	TTGATAATATAAAGTAAATGAAGGAAGTACTGTGTGCAAGAGGGCTATATCTTTTTCCG
SEQ2308	TTGATAATATAAAGCAAATGAAGGAAGTACTGGGTATAAGAGGGCTATATCTTTTTTCCG
SEQ2309	TTGATAATATAAAGCAAATGAAGGAAGTACTGGGTATAAGAGGGCTATATCTTTTTTCCG
SEQ2310	TTGATAATATAAAGTAAATGAAGGAAGTACTGTGCAAGAGGGCTATATCTTTTTCCG
SEQ2311	TTGATAATATAAAGCAAATGAAGGAAGTACTGTGTGCAAGAGGGCTATATCTTTTTTCCG
SEQ2301	GCCCTGTGGGGAGTGGTAAAACAACTCTCATGTATCAATTAGCTTCAGAAGTATTTAAAA
SEQ2302	GCCCTGTGGGGAGTGGTAAAACAACTCTCATGTATCAATTAGCTTCAGAAGTATTTAAAA
SEQ2303	GCCCTGTGGGGAGTGGTAAAACACTCTCATGTATCAATTAGCTTCAGAAGTATTTAAAA
SEQ2304	GCCCTGTGGGGAGTGGTAAAACAACTCTCATGTATCAATTAGCTTCAGAAGTATTTAAAA
SEQ2305	GCCCTGTGGGGAGTGGTAAAACAACTCTCATGTATCAATTAGCTTCAGAAGTATTTAAAA
SEQ2306	GCCCTGTGGGGAGTGGTAAAACAACTCTCATGTATCAATTAGCTTCAGAAGTATTTAAAA GCCCTGTGGGGAGTGGTAAAACAACTCTCATGTATCAATTAGCTTCAGAAGTATTTAAAA
SEQ2307	GCCCTGTGGGGAGTGGTAAAACAACTCTCATGTATCAATTAGCTTCAGAAGTATTTAAAA GCCCTGTGGGGAGTGGTAAAACAACTCTCATGTATCAATTAGCTTCAGAAGTATTTAAAA
SEQ2308	GCCCTGTGGGGAGTGGTAAAACAACTCTCATGTATCAATTAGCTTCAGAAGTATTTAAAA GCCCTGTGGGGAGTGGTAAAACAACTCTCATGTATCAATTAGCTTCAGAAGTATTTAAAA
SEQ2309	GCCCTGTGGGGAGTGGTAAAACAACTCTCATGTATCAATTAGCTTCAGAAGTATTTAAAA GCCCTGTGGGGAGTGGTAAAACAACTCTCATGTATCAATTAGCTTCAGAAGTATTTAAAA
SEQ2310	GCCCTGTGGGGAGTGGTAAAACAACTCTCATGTATCAATTAGCTTCAGAAGTATTTAAAA GCCCTGTGGGGAGTGGTAAAACAACTCTCATGTATCAATTAGCTTCAGAAGTATTTAAAA
SEQ2311	GCCCTGTGGGGGGTGGTAAAACACTCTCATGTATCAAATTAGCTTCAGAAGTATTTAAAA



# Table 23: Comparative Sequences relating to SAG0163 (competence protein Cg1A)

SEQ2301	ATAAGCAAATTATCACGATTGAAGATCCGGTAGAAATCAAGAATGACAAGATGTTACAAC
SEQ2302	ATAAGCAAATTATCACGATTGAAGATCCGGTAGAAATCAAGAATGACAAGATGTTACAAC
SEQ2303	ATAAGCAAATTATCACGATTGAAGATCCGGTAGAAATCAAGAATGACAAGATGTTACAAC
SEQ2304	ATAAGCAAATTATCACGATTGAAGATCCGGTAGAAATCAAGAATGACAAGATGTTACAAC
SEQ2305	ataagcaaattatcacgattgaagatccggtagaaatcaagaatgacaagatgttacaac
SEQ2306	ATAAGCAAATTATCACGATTGAAGATCCGGTAGAAATCAAGAATGACAAGATGTTACAAC
SEQ2307	ataagcaaattatcacgattgaagatccggtagaaatcaagaatgacaagatgttacaac
SEQ2308	ataagcaaattatcacgattgaagatccggtagaaatcaagaatgacaagatgttacaac
SEQ2309	ataagcaaattatcacgattgaagatccggtagaaatcaagaatgacaagatgttacaac
SEQ2310	ataagcaaattatcacgattgaagatccggtagaaatcaagaatgacaagatgttacaac
SEQ2311	ATAAGCAAATTATCACGATTGAAGATCCGGTAGAAATCAAGAATGACAAGATGTTACAAC
SEQ2301	TCCAATTGAATGAGGATATTGGAATGACTTATGATGCTTTAATCAAACTGTCTTTACGGC
SEQ2302	TCCAATTGAATGAGGATATTGGAATGACTTATGATGCTTTAATCAAACTGTCTTTACGGC
SEQ2303	TCCAATTGAATGAGGATATTGGAATGACTTATGATGCTTTAATCAAACTGTCTTTACGGC
SEQ2304	TCCAATTGAATGAGGATATTGGAATGACTTATGATGCTTTAATCAAACTGTCTTTACGGC
SEQ2305	TCCAATTGAATGAGGATATTGGAATGACTTATGATGCTTTAATCAAACTGTCTTTACGGC
SEQ2306	TCCAATTGAATGAGGATATTGGAATGACTTATGATGCTTTAATCAAACTGTCTTTACGGC
SEQ2307	TCCAATTGAATGAGGATATTGGAATGACTTATGATGCTTTAATCAAACTGTCTTTACGGC
SEQ2308	TCCAATTGAATGAGGATATTGGAATGACTTATGATGCTTTAATCAAACTGTCTTTACGGC
SEQ2309	TCCAATTGAATGAGGATATTGGAATGACTTATGATGCTTTAATCAAACTGTCTTTACGGC
SEQ2310	TCCAATTGAATGAGGATATTGGAATGACTTATGATGCTTTAATCAAACTGTCTTTACGGC
SEQ2311	TCCAATTGAATGAGGATATTGGAATGACTTATGATGCTTTAATCAAACTGTCTTTACGGC
SEQ2301	ATCGTCCAGATATTTTAATTATCGGAGAGAT-TAGAGATCAAGCGACGGCCCGTGCTGTT
SEQ2302	ATCGTCCAGATATTTTAATTATCGGAGAGAT-TAGAGATCAAGCGACGGCTCGTGCTGTT
SEQ2303	ATCGTCCAGATATTTTAATTATCGGAGAGAT-TAGAGATCAAGCGACGGCCCGTGCTGTT
SEQ2304	ATCGTCCAGATATTTTAATTATCGGAGAGAT-TAGAGATCAAGCGACGGCCCGTGCTGTT
SEQ2305	ATCGTCCAGATATTTTAATTATCGGAGAGAT-TAGAGATCAAGCGACGCCCGTGCTGTT
SEQ2306	ATCGTCCAGATATTTTAATTATCGGAGAGAT-TAGAGATCAAGCGACGGCCCGTGCTGTT
SEQ2307	ATCGTCCAGATATTTTAATTATCGGAGAGAT-TAGAGATCAAGCGACGGCCCGTGCTGTT
SEQ2308	ATCGTCCAGATATTTTAATTATCGGAGAGAAATAGAGATCAAGCGACGGCCCGTGCTGTT
SEQ2309	ATCGTCCAGATATTTTAATTATCGGAGAGAT-TAGAGATCAAGCGACGGCCCGTGCTGTT
SEQ2310	ATCGTCCAGATATTTTAATTATCGGAGAGAT-TAGAGATCAAGCGACGGCCCGTGCTGTT
SEQ2311	ATCGTCCAGATATTTTAATTATCGGAGAGAT-TAGAGATCAAGCGACGGCCCGTGCTGTT
SEQ2301	ATTCGTGCAAGTTTAACGGGAGTGATGGTTTTTTCTACTATTCATGCTAAAAGTATTTCC
SEQ2302	ATTCGTGCAAGTTTAACGGGAGTGATGGTTTTTTCTACTATTCATGCTAAAAGTATTCCC
SEQ2303	ATTCGTGCAAGTTTAACGGGAGTGATGGTTTTTTCTACTATTCATGCTAAAAGTATTCCC
SEQ2304	ATTCGTGCAAGTTTAACGGGAGTGATGGTTTTTTCTACTATTCATGCTAAAAGTATTCCC
SEQ2305	ATTCGTGCAAGTTTAACGGGAGTGATGGTTTTTTCTACTATTCATGCTAAAAGTATTCCC
SEQ2306	ATTCGTGCAAGTTTAACGGGAGTGATGGTTTTTTCTACTATTCATGCTAAAAGTATTTCC
SEQ2307	ATTCGTGCAAGTTTAACGGGAGTAATGGTTTTTTCTACTATTCATGCTAAAAGTATTCCC
SEQ2308	ATTCGTGCAAGTTTAACGGGAGTGATGTTTTTTTCTACTATTCATGCTAAAAGTATTCCC
SEQ2309	ATTCGTGCAAGTTTAACGGGAGTGATGGTTTTTTCTACTATTCATGCTAAAAGTATTCCC
SEQ2310 SEQ2311	ATTCGTGCAAGTTTAACGGGAGTAATGGTTTTTTCTACTATTCATGCTAAAAGTATTCCC ATTCGTGCAAGTTTAACGGGAGTAATGGTTTTTCTACTATTCATGCTAAAAGTATTCCC
SEQ2301	GGAGTCTATGATAGGCTTATAGAATTAGGGGTTAACTATCAAGAGTTAGAAAATAGTCTA
SEQ2302	GGAGTCTATGATAGGCTTATAGAATTAGGGGTTAACTATCAAGAGTTAGAAAATAGTCTA
SEQ2303	GGAGTCTATGATAGGCTTATAGAATTAGGGGTTAACTATCAAGAGTTAGAAAATAGTCTA
SEQ2304	GGAGTCTATGATAGGCTTATAGAATTAGGGGTTAACTATCAAGAGTTAGAAAATAGTCTA
SEQ2305	GGAGTCTATGATAGGCTTATAGAATTAGGGGTTAACTATCAAGAGTTAGAAAATAGTCTA
SEQ2306	GGAGTCTATGATAGGCTTATAGAATTAGGGGTTAACTATCAAGAGTTAGAAAATAGTCTA
SEQ2307 SEQ2308	GGAGTCTATGATAGGCTTATAGAATTAGGGGTTAACTATCAAGAGTTAGAAAATAGTCTA
SEQ2309	GGAGTCTATGATAGGCTTATAGAATTAGGGGTTAACTATCAAGAGTTAGAAAATAGTCTA
SEQ2309 SEQ2310	GGAGTCTATGATAGGCTTATAGAATTAGGGGTTAACTATCAAGAGTTAGAAAATAGTCTA GGAGTCTATGATAGGCTTATAGAATTAGGGGTTAACTATCAAGAGTTAGAAAATAGTCTA
SEQ2311	GGAGTCTATGATAGGCTTATAGAATTAGGGGTTAACTATCAAGAGTTAGAAAATAGTCTA
SEQ2301	ሽ ሽ ሽመመሽ ሽመሽ ሮሮሽ መስመሮሽ ሽ ድርቁመው ሽ ሽመውረስ እንደረደ ሽ ሽ ድርድመሽ ሽ መመረስ ድመመውር ችር እና እና ድርመ
SEQ2301 SEQ2302	AAATTAATAGCATATCAACGTTTAATTGGAGGAGGAAGCCTAATTGACTTTGAGACAGGT  AAATTAATAGCATATCAACGTTTAATTGGAGGAGGAAGCCTAATTGACTTTGAGACAACAACAACAAAACTAAATTGACAACAACAACAAACTTTTGAGACAACAACAACAACAAACA
SEQ2302 SEQ2303	AAATTAATAGCATATCAACGTTTAATTGGAGGAGGAAGCCTAATTGACTTTGAGACAAGT AAATTAATAGCATATCAACGTTTAATTGGAGGAGGAAGCCTAATTGACTTTGAGACAGGT
SEQ2304	AAATTAATAGCATATCAACGTTTAATTGGAGGAGGAAGCCTAATTGACTTTGAGACAGGT AAATTAATAGCATATCAACGTTTAATTGGAGGAGGAAGCCTAATTGACTTTGAGACAGGT
SEQ2305	AAATTAATAGCATATCAACGTTTAATTGGAGGAGGAAGCCTAATTGACTTTGAGACAGGT AAATTAATAGCATATCAACGTTTAATTGGAGGAGGAAGCCTAATTGACTTTGAGACAGGT
SEQ2306	AAATTAATAGCATATCAACGTTTAATTGGAGGAGGAAGCCTAATTGACTTTGAGACAGGT
SEQ2307	AAATTAATAGCATATCAACGTTTAATTGGAGGAGGAAGCCTAATTGACTTTGAGACAAGT





SEQ2308	AAATTAATAGCATATCAACGTTTAATTGGAGGAGGAAGCCTAATTGACTTTGAGACAGGT
SEQ2309	AAATTAATAGCATATCAACGTTTAATTGGAGGAGGAAGCCTAATTGACTTTGAGACAGGT
SEQ2310	AAATTAATAGCATATCAACGTTTAATTGGAGGAGGAAGCCTAATTGACTTTGAGACAAGT
SEQ2311	AAATTAATAGCATATCAACGTTTAATTGGAGGAGGAAGCCTAATTGACTTTGAGACAAGT
SEQ2301	AACTTTAAAAAACACTCATCAGACAAGTGGAATAGACAAGTGGATATCTTGGCTGAAGAA
SEQ2302	AACTTTAAAAAACACTCATCAGACAAGTGGAATAGACAAGTGGATATCTTGGCTGAAGAA
SEQ2303	AATTTTAAAAAACACTCATCAGACAAGTGGAATAGACAAGTGGATATCTTGGCTGAAGAA
SEQ2304	AATTTTAAAAAACACTCATCAGACAAGTGGAATAGACAAGTGGATATCTTGGCTGAAGAA
SEQ2305	AATTTTAAAAAACACTCATCAGACAAGTGGAATAGACAAGTGGATATCTTGGCTGAAGAA
SEQ2306	AACTTTAAAAAACACTCATCAGACAAGTGGAATAGACAAGTGGATATCTTGGCTGAAGAA
SEQ2307	AACTTTAAAAAACACTCATCAGACAAGTGGAATAGACAAGTGGATATCTTGGCTGAAGAA
SEQ2308	AATTTTAAAAAACACTCATCAGACAAGTGGAATAGACAAGTGGATATCTTGGCTGAAGAA
SEQ2309	AATTTTAAAAAACACTCATCAGACAAGTGGAATAGACAAGTGGATATCTTGGCTGAAGAA
SEQ2310	AACTTTAAAAAACACTCATCAGACAAGTGGAATAGACAAGTGGATATCTTGGCTGAAGAA
SEQ2311	AACTTTAAAAAACACTCATCAGACAAGTGGAATAGACAAGTGGATATCTTGGCTGAAGAA
SEQ2301	GGACATATCAGTAAGAAACAGGCACAAGT-CGAAAAAATTATCCCTCAAGAAACAACGGA
SEQ2302	GGATATATCAGTAAGAAACAGGCACAAGT-CGAAAAAATTATCCCTCAAGAAACAACGGA
SEQ2303	GGACATATCAGTAAGAAACAGGCACAAGT-CGAAAAAATTATCCCTCAAGAAACAACGGA
SEQ2304	GGACATATCAGTAAGAAACAGGCACAAGTGCGAAAAAATTATCCCTCAAGAAACAACGGA
SEQ2305	GGACATATCAGTAAGAAACAGGCACAAGT-CGAAAAAATTATCCCTCAAGAAACAACGGA
SEQ2306	GGACATATCAGTAAGAAACAGGCACAAGT-CGAAAAAATTATCCCTCAAGAAACAACGGA
SEQ2307	GGACATATCAGTAAGAAACAGGCACAAGT-CGAAAAAATTATCCCTCAAGAAACAACGGA
SEQ2308	GGACATATCAGTAAGAAACAGGCACAAGT-CGAAAAAATTATCCCTCAAGAAACAACGGA
SEQ2309	GGACATATCAGTAAGAAACAGGCACAAGT-CGAAAAAATTATCCCTCAAGAAACAACGGA
SEQ2310	GGACATATCAGTAAGAAACAGGCACAAGT-CGAAAAAATTATCCCTCAAGAAACAACGGA
SEQ2311	GGACATATCAGTAAGAAACAGGCACAAGT-CGAAAAAATTATCCCTCAAGAAACAACGGA
000000	
SEQ2301	AAGTAGTCCAACTTTTAAGTAGTCCAACTTTT
SEQ2302 SEQ2303	AAGTAGTCCAACTTTTAAGTAGTCCAACTTTT
SEQ2303 SEQ2304	AAGTAGTCCAACTTTT
SEQ2304 SEO2305	AAGTAGTCCAACTTTTAAGTAGTCCAACTTTT
SEQ2305 SEQ2306	AAGTAGTCCAACTTTT
SEQ2307	AAGTAGTCCAACTTTTAAGTAGTCCAACTTTT
SEQ2307 SEQ2308	AAGTAGTCCAACTTTTAAGTAGTCCAACTTTT
SEQ2309	AAGTAGTCCAACTTTTAAGTAGTCCAACTTTT
SEQ2310	AAGTAGTCCAACTTTT
SEQ2311	AAGTAGTCCAACTTTT
OUGESTY	ANGINGI CONNCI I I I

#### >SEQ ID NO 6350:63 090 frame: 2

AVEVNAQDIYIIPKGDCYELYMRIDDERRFIDVFEFNRMASLISHFKFVAGMNVGEKRRS QLGSCDYELSEGRLVSLRLSSVGDYRGQESLVIRILYSGHQDLKYWFDNIKQMKEVLGTR GLYLFSGPVGSGKTTLMYQLASEVFKNKQIITIEDPVEIKNDKMLQLQLNEDIGMTYDAL IKLSLRHRPDILIIGEIRDQATARAVIRASLTGVMVFSTIHAKSISGVYDRLIELGVNYQ ELENSLKLIAYQRLIGGGSLIDFETGNFKKHSSDKWNRQVDILAEEGHISKKQAQVEKII PQETTESSPTF

#### >SEQ ID NO 6351:63\_1169NT frame: 3

.LL.NLYYCVFDDERRFIDVFEFNRMASLISHFKFVAGMNVGEKRRSQLGSCDYELSEGR LVSLRLSSVGDYRGQESLVIRILYSGHQDLKYWFDNIKQMKEVLGTRGLYLFSGPVGSGK TTLMYQLASEVFKNKQIITIEDPVEIKNDKMLQLQLNEDIGMTYDALIKLSLRHRPDILI IGEIRDQATARAVIRASLTGVMVFSTIHAKSIPGVYDRLIELGVNYQELENSLKLIAYQR LIGGGSLIDFETSNFKKHSSDKWNRQVDILAEEGYISKKQAQVEKIIPQETTESSPTF

#### >SEQ ID NO 6352:63\_18R821 frame: 1

VQSLAKQVIHQAVEVNAQDIYIIPKGDCYELYMRIDDERRFIDVFEFNRMASLISHFKFV AGMNVGEKRRSQLGSCDYELSEGRLVSLRLSSVGDYRGQESLVIRILYSGHQDLKYWFDN IKQMKEVLGIRGLYLFSGPVGSGKTTLMYQLASEVFKNKQIITIEDPVEIKNDKMLQLQL NEDIGMTYDALIKLSLRHRPDILIIGEIRDQATARAVIRASLTGVMVFSTIHAKSIPGVY DRLIELGVNYQELENSLKLIAYQRLIGGGSLIDFETGNFKKHSSDKWNRQVDILAEEGHI SKKQAQVEKIIPQETTESSPTF

#### >SEQ ID NO 6353: 63 2603 frame: 1

DIYIIPKGDCYELYMRIDDERRFIDVFEFNRMASLISHFKFVAGMNVGEKRRSQLGSCDY ELSEGRLVSLRLSSVGDYRGQESLVIRILYSGHQDLKYWFDNIKOMKEVLGIRGLYLFSG



## Table 23: Comparative Sequences relating to SAG0163 (competence protein Cg1A)

PVGSGKTTLMYQLASEVFKNKQIITIEDPVEIKNDKMLQLQLNEDIGMTYDALIKLSLRH RPDILIIGEIRDQATARAVIRASLTGVMVFSTIHAKSIPGVYDRLIELGVNYQELENSLK LIAYQRLIGGGSLIDFETGNFKKHSSDKWNRQVDILAEEGHISKKQAQVRKNYPSRNNGK .SNF

#### >SEQ ID NO 6354:63 A909 frame: 1

VQSLAKQVIHQAVEVNAQDIYIIPKGDCYELYMRIDDERRFIDVFEFNRMASLISHFKFV AGMNVGEKRRSQLGSCDYELSEGRLVSLRLSSVGDYRGQESLVIRILYSGHQDLKYWFDN IKQMKEVLGIRGLYLFSGPVGSGKTTLMYQLASEVFKNKQIITIEDPVEIKNDKMLQLQL NEDIGMTYDALIKLSLRHRPDILIIGEIRDQATARAVIRASLTGVMVFSTIHAKSIPGVY DRLIELGVNYQELENSLKLIAYQRLIGGGSLIDFETGNFKKHSSDKWNRQVDILAEEGHI SKKQAQVEKIIPQETTESSPTF

#### >SEQ ID NO 6355:63 CJB110 frame: 1

VQSLAKQVIHQAVEVNAQDIYIIPKGDCYELYMRIDDERRFIDVFEFNRMASLISHFKFV AGMNVGEKRRSQLGSCDYELSEGRLVSLRLSSVGDYRGQESLVIRILYSGHQDLKYWFDN IKQMKEVLGTRGLYLFSGPVGSGKTTLMYQLASEVFKNKQIITIEDPVEIKNDKMLQLQL NEDIGMTYDALIKLSLRHRPDILIIGEIRDQATARAVIRASLTGVMVFSTIHAKSISGVY DRLIELGVNYQELENSLKLIAYQRLIGGGSLIDFETGNFKKHSSDKWNRQVDILAEEGHI SKKQAQVEKIIPQETTESSPTF

#### >SEQ ID NO 6356:63 CJB110 frame: 1

VQSLAKQVIHQAVEVNAQDIYIIPKGDCYELYMRIDDERRFIDVFEFNRMASLISHFKFV AGMNVGEKRRSQLGSCDYELSEGRLVSLRLSSVGDYRGQESLVIRILYSGHQDLKYWFDN IKQMKEVLGTRGLYLFSGPVGSGKTTLMYQLASEVFKNKQIITIEDPVEIKNDKMLQLQL NEDIGMTYDALIKLSLRHRPDILIIGEIRDQATARAVIRASLTGVMVFSTIHAKSISGVY DRLIELGVNYQELENSLKLIAYQRLIGGGSLIDFETGNFKKHSSDKWNRQVDILAEEGHI SKKQAQVEKIIPQETTESSPTF

#### >SEQ ID NO 6357: 63\_H36B frame: 1

SLAKQVIHQAVEVNAQDIYIIPKGDCYELYMRIDDERRFIDVFEFNRMASLISHFKFVAG MNVGEKRRSQLGSCDYELSEGRLVSLRLSSVGDYRGQESLVIRILYSGHQDLKYWFDNIK QMKEVLGIRGLYLFSGPVGSGKTTLMYQLASEVFKNKQIITIEDPVEIKNDKMLQLQLNE DIGMTYDALIKLSLRHRPDILIIGEK

#### >SEQ ID NO 6358:63\_JM9130013 frame: 1

VQSLAKQVIHQAVEVNAQDIYIIPKGDCYELYMRIDDERRFIDVFEFNRMASLISHFKFV AGMNVGEKRRSQLGSCDYELSEGRLVSLRLSSVGDYRGQESLVIRILYSGHQDLKYWFDN IKQMKEVLGIRGLYLFSGPVGSGKTTLMYQLASEVFKNKQIITIEDPVEIKNDKMLQLQL NEDIGMTYDALIKLSLRHRPDILIIGEIRDQATARAVIRASLTGVMVFSTIHAKSIPGVY DRLIELGVNYQELENSLKLIAYQRLIGGGSLIDFETGNFKKHSSDKWNRQVDILAEEGHI SKKQAQVEKIIPQETTESSPTF

#### >SEQ ID NO 6359:63\_M732 frame: 3

TCYETLYAYLMMKRRFIDVFEFNRMASLISHFKFVAGMNVGEKRRSQLGSCDYELSEGRL VSLRLSSVGDYRGQESLVIRTLYSGHQDLKYWFDNIK.MKEVLCARGLYLFSGPVGSGKT TLMYQLASEVFKNKQIITIEDPVEIKNDKMLQLQLNEDIGMTYDALIKLSLRHRPDILII GEIRDQATARAVIRASLTGVMVFSTIHAKSIPGVYDRLIELGVNYQELENSLKLIAYQRL IGGGSLIDFETSNFKKHSSDKWNRQVDILAEEGHISKKQAQVEKIIPQETTESSPTF >SEQ ID NO 6360:63 M781 frame: 3

VEVNAQDIYIIPKGDCYEFYMRIDDERRFIDVFEFNRMASLISHFKFVAGMNVGEKRRSQ LGSCDYELSEGRLVSLRLSSVGDYRGQESLVIRTLYSGHQDLKYWFDNIKQMKEVLCARG LYLFSGPVGSGKTTLMYQLASEVFKNKQIITIEDPVEIKNDKMLQLQLNEDIGMTYDALI KLSLRHRPDILIIGEIRDQATARAVIRASLTGVMVFSTIHAKSIPGVYDRLIELGVNYQE LENSLKLIAYQRLIGGGSLIDFETSNFKKHSSDKWNRQVDILAEEGHISKKQAQVEKIIP QETTESSPTF

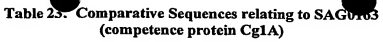
#### >SEQ ID NO 6361:63\_COH1 frame: 3

VIVMKFYMRIDDERRFIDVFEFNRMASLISHFKFVAGMNVGEKRRSQLGSCDYELSEGRL VSLRLSSVGDYRGQESLVIRTLYSGHQDLKYWFDNIK



# Table 23: Comparative Sequences relating t SAG0163 (competence protein Cg1A)

SEQ6350	AVEVNAQDIYIIPKGDCYELYMRIDDERRFIDVFEFNRMASLISHFKFV
SEQ6351	LINLYYCVFDDERRFIDVFEFNRMASLISHFKFV
SEQ6352	QSLAKQVIHQAVEVNAQDIYIIPKGDCYELYMRIDDERRFIDVFEFNRMASLISHFKFV
SEQ6353	DIYIIPKGDCYELYMRIDDERRFIDVFEFNRMASLISHFKFV
SEQ6354	QSLAKQVIHQAVEVNAQDIYIIPKGDCYELYMRIDDERRFIDVFEFNRMASLISHFKFV
SEQ6355	QSLAKQVIHQAVEVNAQDIYIIPKGDCYELYMRIDDERRFIDVFEFNRMASLISHFKFV
SEQ6356	QSLAKQVIHQAVEVNAQDIYIIPKGDCYELYMRIDDERRFIDVFEFNRMASLISHFKFV
SEQ6357	-SLAKQVIHQAVEVNAQDIYIIPKGDCYELYMRIDDERRFIDVFEFNRMASLISHFKFV
SEQ6358	QSLAKQVIHQAVEVNAQDIYIIPKGDCYELYMRIDDERRFIDVFEFNRMASLISHFKFV
SEQ6359	typetlyaylmmkrrfidvfefnrmaslishfkfv
SEQ6360	vevnaqdiyiipkgdcyefymridderrfidvfefnrmaslishfkfv
SEQ6361	VIVMKFYMRIDDERRFIDVFEFNRMASLISHFKFV
SEQ6350	AGMNVGEKRRSQLGSCDYELSEGRLVSLRLSSVGDYRGQESLVIRILYSGHQDLKYWFDN
SEQ6351	AGMNVGEKRRSQLGSCDYELSEGRLVSLRLSSVGDYRGQESLVIRILYSGHQDLKYWFDN
SEQ6352	AGMNVGEKRRSQLGSCDYELSEGRLVSLRLSSVGDYRGQESLVIRILYSGHQDLKYWFDN
SEQ6353	AGMNVGEKRRSQLGSCDYELSEGRLVSLRLSSVGDYRGQESLVIRILYSGHQDLKYWFDN
SEQ6354	AGMNVGEKRRSQLGSCDYELSEGRLVSLRLSSVGDYRGQESLVIRILYSGHQDLKYWFDN
SEQ6355	AGMNVGEKRRSQLGSCDYELSEGRLVSLRLSSVGDYRGQESLVIRILYSGHQDLKYWFDN
SEQ6356	AGMNVGEKRRSQLGSCDYELSEGRLVSLRLSSVGDYRGQESLVIRILYSGHQDLKYWFDN
SEQ6357	AGMNVGEKRRSQLGSCDYELSEGRLVSLRLSSVGDYRGQESLVIRILYSGHQDLKYWFDN
SEQ6358	AGMNVGEKRRSQLGSCDYELSEGRLVSLRLSSVGDYRGQESLVIRILYSGHQDLKYWFDN
SEQ6359	AGMNVGEKRRSQLGSCDYELSEGRLVSLRLSSVGDYRGQESLVIRTLYSGHQDLKYWFDN
SEQ6360	AGMNVGEKRRSQLGSCDYELSEGRLVSLRLSSVGDYRGQESLVIRTLYSGHQDLKYWFDN
SEQ6361	AGMNVGEKRRSQLGSCDYELSEGRLVSLRLSSVGDYRGQESLVIRTLYSGHQDLKYWFDN
SEQ6350	IKOMKEVLGTRGLYLFSGPVGSGKTTLMYQLASEVFKNKQIITIEDPVEIKNDKMLQLQL
SEQ6351	IKOMKEVLGTRGLYLFSGPVGSGKTTLMYOLASEVFKNKOIITIEDPVEIKNDKMLOLOL
SEQ6352	IKOMKEVLGIRGLYLFSGPVGSGKTTLMYQLASEVFKNKQIITIEDPVEIKNDKMLQLQL
SEQ6353	IKOMKEVLGIRGLYLFSGPVGSGKTTLMYQLASEVFKNKQIITIEDPVEIKNDKMLQLQL
SEQ6354	IKOMKEVLGIRGLYLFSGPVGSGKTTLMYOLASEVFKNKQIITIEDPVEIKNDKMLQLQL
SEQ6355	IKOMKEVLGTRGLYLFSGPVGSGKTTLMYQLASEVFKNKQIITIEDPVEIKNDKMLQLQL
SEQ6356	IKOMKEVLGTRGLYLFSGPVGSGKTTLMYQLASEVFKNKQIITIEDPVEIKNDKMLQLQL
SEQ6357	IKOMKEVLGIRGLYLFSGPVGSGKTTLMYQLASEVFKNKQIITIEDPVEIKNDKMLQLQL
SEQ6358	IKOMKEVLGIRGLYLFSGPVGSGKTTLMYQLASEVFKNKQIITIEDPVEIKNDKMLQLQL
SEQ6359	IK-MKEVLCARGLYLFSGPVGSGKTTLMYQLASEVFKNKQIITIEDPVEIKNDKMLQLQL
SEQ6360	IKQMKEVLCARGLYLFSGPVGSGKTTLMYQLASEVFKNKQIITIEDPVEIKNDKMLQLQL
SEQ6361	IK
SEQ6350	EDIGMTYDALIKLSLRHRPDILIIGEIRDQATARAVIRASLTGVMVFSTIHAKSISGVY
SEQ6351	EDIGMTYDALIKLSLRHRPDILIIGEIRDOATARAVIRASLTGVMVFSTIHAKSIPGVY
SEQ6352	EDIGMTYDALIKLSLRHRPDILIIGEIRDQATARAVIRASLTGVMVFSTIHAKSIPGVY
SEQ6353	EDIGMTYDALIKLSLRHRPDILIIGEIRDOATARAVIRASLTGVMVFSTIHAKSIPGVY
SEQ6354	EDIGMTYDALIKLSLRHRPDILIIGEIRDOATARAVIRASLTGVMVFSTIHAKSIPGVY
SEQ6355	EDIGMTYDALIKLSLRHRPDILIIGEIRDOATARAVIRASLTGVMVFSTIHAKSISGVY
SEQ6356	EDIGMTYDALIKLSLRHRPDILIIGEIRDQATARAVIRASLTGVMVFSTIHAKSISGVY
SEQ6357	EDIGMTYDALIKLSLRHRPDILIIGEK
SEQ6358	EDIGMTYDALIKLSLRHRPDILIIGEIRDQATARAVIRASLTGVMVFSTIHAKSIPGVY
SEQ6359	EDIGMTYDALIKLSLRHRPDILIIGEIRDQATARAVIRASLTGVMVFSTIHAKSIPGVY
SEQ6360	EDIGMTYDALIKLSLRHRPDILIIGEIRDQATARAVIRASLTGVMVFSTIHAKSIPGVY
SEQ6361	
SEQ6350	RLIELGVNYQELENSLKLIAYQRLIGGGSLIDFETGNFKKHSSDKWNRQVDILAEEGHI
SEQ6351	RLIELGVNYQELENSLKLIAYQRLIGGGSLIDFETSNFKKHSSDKWNRQVDILAEEGYI
SEQ6352	RLIELGVNYQELENSLKLIAYQRLIGGGSLIDFETGNFKKHSSDKWNRQVDILAEEGHI
SEQ6353	RLIELGVNYQELENSLKLIAYQRLIGGGSLIDFETGNFKKHSSDKWNRQVDILAEEGHI
SEQ6354	RLIELGVNYQELENSLKLIAYQRLIGGGSLIDFETGNFKKHSSDKWNRQVDILAEEGHI
SEQ6355	RLIELGVNYQELENSLKLIAYQRLIGGGSLIDFETGNFKKHSSDKWNRQVDILAEEGHI
SEQ6356	RLIELGVNYQELENSLKLIAYQRLIGGGSLIDFETGNFKKHSSDKWNRQVDILAEEGHI
SEQ6357	
SEQ6358	RLIELGVNYQELENSLKLIAYQRLIGGGSLIDFETGNFKKHSSDKWNRQVDILAEEGHI
SEQ6359	RLIELGVNYQELENSLKLIAYQRLIGGGSLIDFETSNFKKHSSDKWNRQVDILAEEGHI
SEQ6360	RLIELGVNYQELENSLKLIAYQRLIGGGSLIDFETSNFKKHSSDKWNRQVDILAEEGHI
SEQ6361	



SEQ6350	KKQAQVEKIIPQETTESSPTF
SEQ6351	KKQAQVEKIIPQETTESSPTF
SEQ6352	KKQAQVEKIIPQETTESSPTF
SEQ6353	KKQAQVRKNYPSRNNGKSNF-
SEQ6354	KKQAQVEKIIPQETTESSPTF
SEQ6355	KKQAQVEKIIPQETTESSPTF
SEQ6356	KKQAQVEKIIPQETTESSPTF
SEQ6357	
SEQ6358	KKQAQVEKIIPQETTESSPTF
SEQ6359	KKQAQVEKIIPQETTESSPTF
SEQ6360	KKQAQVEKIIPQETTESSPTF
SEQ6361	

## Figure 24: Comparative Sequences relating to SAG0290 (ABC transporter, substrate-binding protein)

## SEQ ID NO. 2101: SAG0290 FROM THE 1169NT1 GBS TYPE V STRAIN (REVERSE COMPLEMENT)

### SEQ ID NO. 2402: SAG0290 FROM THE 18RS21 GBS TYPE II STRAIN (REVERSE COMPLEMENT)

### SEQ ID NO. 2403: SAG0290 FROM THE 2603 V/R GBS TYPE V STRAIN (REVERSE COMPLEMENT)



## Figure 24: Comparative Sequences relating to SAG0290 (ABC transporter, substrate-binding protein)

## SEQ ID NO. 2405: SAG0290 FROM THE A909 GBS TYPE Ia STRAIN (REVERSE COMPLEMENT)

GTATCAGTTCAGGCGTCAGAGAAAGTAGAACTTAAAGTAGCTACAGATTCTGACACGGCACCATTTACTTA
TCAAAAAGACGGGAAATTCAAAGGTTATGATGTTGATGTTGTCAAAGCTGTTTTTTAAAGGTAGTAACA
AAGTAACCTTCAAGACAGTTCCTTTTGATACTATTTCAACAGGTATTGATGCAGGGAAATTTGATTTATCA
GCTAATGATTTTTCATACAATAAAGAAAGAGCAGAAAAATATCTCTTCTCAGATCCTATATCCCGTTCAAA
TTATGCCGTAGTAGGGAAGAAGGGGAGCCATTACAAATCATTAAGTGACCTCTCTGGAAAATCAACCGAAG
TTTTATCTGGCGTTAACTATGCACAGGTTCTAGAAAATTGGAATAAAAATCATNNTAATAAAAAACCANTA
AAAATNAAATATGTTTCTGGGACAACTGGTGTTACTAGCAGATTAAAAAATATTGAGAGTGGGAAAATTGA
CTTTATCCTATATGATGCCATTTCATCCGACTATATTGTAAAAGACCAATCATTAAACTTAAGCGTTTCTC
CTTTGAAAGGTAAAATTGGTAATAATAAGGATGGACTAGAATACCTCCTTTTTACCAAAAGATAAAAAAGGT
AAAACTCTACAGAAATTTATAAATAAGCGT

### SEQ ID NO. 2406: SAG0290 FROM THE CJB110 GBS NONTYPEABLE STRAIN (REVERSE COMPLEMENT)

### SEQ ID NO. 2407: SAG0290 FROM THE COH1 GBS TYPE III STRAIN (REVERSE COMPLEMENT)

### SEQ ID NO. 2408: SAG0290 FROM THE H36b GBS TYPE ID STRAIN (REVERSE COMPLEMENT)

## Figur 24: Comparative Sequences relating to SAG0290 (ABC transp rter, substrate-binding protein)

## SEQ ID NO. 2409: SAG0290 FROM THE JM9130013 GBS STRAIN VIII (REVERSE COMPLEMENT)

## SEQ ID NO. 2410: SAG0290 FROM THE M732 GBS TYPE III STRAIN (REVERSE COMPLEMENT)

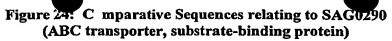
## SEQ ID NO. 2411: SAG0290 FROM THE M781 GBS TYPE III STRAIN (REVERSE COMPLEMENT)

SEQ2401	TATCAGTTCAGGCGTCAGAGAAAGTAGAACTTAAAGTAGCTACAGATTCTGACACGGCA
SEQ2402 SEQ2403	TATCAGTTCAGGCGTCAGAGAAAGTAGAACTTAAAGTAGCTACAGATTCTGACACGGCA
SEQ2404	TATCAGTTCAGGCGTCAGAGAAAGTAGAACTTAAAGTAGCTACAGATTCTGACACGGCA
SEQ2405	TATCAGTTCAGGCGTCAGAGAAAGTAGAACTTAAAGTAGCTACAGATTCTGACACGGCA
SEQ2406	TATCAGTTCAGGCGTCAGAGAAAGTAGAACTTAAAGTAGCTACAGATTCTGACACGGCA
SEQ2407	TATCAGTTCAGGCGTCAGAGAAAGTAGAACTTAAAGTAGCTACAGATTCTGACACGCCA
SEQ2408	TATCAGTTCAGGCGTCAGAGAAAGTAGAACTTAAAGTAGCTACAGATTCTGACACGGCA
SEQ2409	TATCAGTTCAGGCGTCAGAGAAAGTAGAACTTAAAGTAGCTACAGATTCTGACACGGCA
SEQ2410	TATCAGTTCAGGCGTCAGAGAAAGTAGAACTTAAAGTAGCTACAGATTCTGACACGGCA
SEQ2411	TATCAGTTCAGGCGTCAGAGAAAGTAGAACTTAAAGTAGCTACAGATTCTGACACGGCA
	•
SEQ2401	CATTTACTTATCAAAAAGACGGGAAATTCAAAGGTTATGATGTTGATGTTGTCAAAGCT
SEQ2402	CATTTACTTATRAAAAAGACGGGAAATTCAAAGGTTATGATGTTGATGTTGTCAAAGCT
SEQ2403	ATTCAAAGGTTATGATGTTGATGTTGTCAAAGCT
SEQ2404	CATTTACTTATCAAAAAGACGGGAAATTCAAAGGTTATGATGTTGATGTTGTCAAAGCT
SEQ2405	CATTTACTTATCAAAAAGACGGGAAATTCAAAGGTTATGATGTTGATGTTGTCAAAGCT
SEQ2406	CATTTACTTATCAAAAAGACGGGAAATTCAAAGGTTATGATGTTGATGTTGTCAAAGCT
SEQ2407	CATTTACTTATCAAAAAGACGGGAAATTCAAAGGTTATGACGTTGATGTTGTCAAAGCT
SEQ2408	
SEQ2409	CATTTACTTATCAAAAAAGACGGGAAATTCAAAGGTTATGATGTTGATGTTGTCAAAGCT
DUKEAAA	CATTTACTTATCAAAAAGACGGGAAATTCAAAGGTTATGATGTTGATGTTGTCAAAGCT



# Figure 24: Comparative Sequences relating to SAG0290 (ABC transporter, substrate-binding protein)

SEQ2410	Catttacttatcaaaaagacgggaaattcaaaggttatgacgttgatgttgtcaaagct
SEQ2411	Catttacttatcaaaagacggaaattcaaaggttatgacgttgatgttgtcaaagct
SEQ2401	GTTTTTAAAGGTAGTAAGTACAAAGTAACCTTCAAGACAGTTCCTTTTGATACTATTTCA
SEQ2402	GTTTTTAAAGGTAGTAAGTACAAAGTAACCTTCAAGACAGTTCCTTTTGATACTATTTCA
SEQ2403	GTTTTTAAAGGTAGTAAGTACAAAGTAACCTTCAAGACAGTTCCTTTTGATACTATTTCA
SEQ2404	GTTTTTAAAGGTAGTAGTACAAAGTAACCTTCAAGACAGTTCCTTTTGATACTATTTCA
SEQ2405	GTTTTTAAAGGTAGTAAGTACAAAGTAACCTTCAAGACAGTTCCTTTTGATACTATTTCA
SEQ2406	GTTTTTAAAGGTAGTAAGTACAAAGTAACCTTCAAGACAGTTCCTTTTGATACTATTTCA
SEQ2407	GTTTTTAAAGGTAGTAAGTACAAAGTAACCTTCAAGACAGTTCCTTTTGATACTATTTCA
SEQ2408	GTTTTTAAAGGTAGTAAGTACAAAGTAACCTTCAAGACAGTTCCTTTTGATACTATTTCA
SEQ2409	GTTTTTAAAGGTAGTAAGTACAAAGTAACCTTCAAGACAGTTCCTTTTGATACTATTTCA
_	
SEQ2410	GTTTTTAAAGGTAGTAAGTACAAAGTAACCTTCAAGACAGTTCCTTTTGATACTATTTCA
SEQ2411	GTTTTTAAAGGTAGTAAGTACAAAGTAACCTTCAAGACAGTTCCTTTTGATACTATTTCA
SEQ2401	ACAGGTATTGATGCAGGGAAATTTGATTTATCAGCTAATGATTTTTCATACAATAAAGAA
SEQ2402	ACAGGTATTGATGCAGGGAAATTTGATTTATCAGCTAATGATTTTTCATACAATAAAGAA
SEQ2403	ACAGGTATTGATGCAGGGAAATTTGATTTATCAGCTAATGATTTTTCATACAATAAAGAA
SEQ2404	ACAGGTATTGATGCAGGGAAATTTGATTTATCAGCTAATGATTTTTCATACAATAAAGAA
SEQ2405	ACAGGTATTGATGCAGGGAAATTTGATTTATCAGCTAATGATTTTTCATACAATAAAGAA
SEQ2406	ACAGGTATTGATGCAGGGAAATTTGATTTATCAGCTAATGATTTTTCATACAATAAAGAA
SEQ2407	ACAGGTATTGATGCAGGGAAATTTGATTATCAGCTAATGATTTTTCATATAAAAGAA
SEQ2408	ACAGGTATTGATGCAGGGAAATTTGATTTATCAGCTAATGATTTTTCATACAATAAAGAA
SEQ2409	ACAGGTATTGATGCAGGGAAATTTGATTTATCAGCTAATGATTTTTCATACAATAAAGAA
SEQ2410	ACAGGTATTGATGCAGGGAAATTTGATTTATCAGCTAATGATTTTTCATATAATAAAGAA
SEQ2411	ACAGGTATTGATGCAGGGAAATTTGATTATCAGCTAATGATTTTTCATATAATAAGAA
SEQ2401	AGAGCAGAAAAATATCTCTTCTCAGACCCTATATCCCGTTCAAATTATGCCGTAGTAGGG
SEQ2402	AGAGCAGAAAAATATCTCTTCTCAGATCCTATATCCCGTTCAAATTATGCCGTAGTAGGG
SEQ2403	AGAGCAGAAAAATATCTCTCCCAGATCCTATATCCCGTTCAAATTATGCCGTAGTAGGG
SEQ2404	AGAGCAGAAAAATATCTCTTCTCAGATCCTATATCCCGTTCAAATTATGCCGTAGTAGGG
SEQ2405	AGAGCAGAAAAATATCTCTTCTCAGATCCTATATCCCGTTCAAATTATGCCGTAGTAGGG
SEQ2405	
<del></del>	AGAGCAGAAAATATCTCTCTCAGATCCTATATCCCGTTCAAATTATGCCGTAGTAGGG
SEQ2407	AGAGCAGAAAAATATCTCTTCTCAGATCCTATATCCCGTTCAAATTATGCCGTAGTAGGG
SEQ2408	AGAGCAGAAAAATATCTCTTCTCAGATCCTATATCCCGTTCAAATTATGCCGTAGTAGGG
SEQ2409	AGAGCAGAAAAATATCTCTCTCAGATCCTATATCCCGTTCAAATTATGCCGTAGTAGGG
SEQ2410	AGAGCAGAAAAATATCTCTTCTCAGATCCTATATCCCGTTCAAATTATGCCGTAGTAGGG
SEQ2411	AGAGCAGAAAATATCTCTTCTCAGATCCTATATCCCGTTCAAATTATGCCGTAGTAGGG
SEQ2401	AAGAAGGGGAGCCATTACAAATCATTAAGTGACCTCTCTGGAAAATCAACAGAAGTTTTA
SEQ2402	AAGAAGGGGAGCCATTACAAATCATTAAGTGACCTCTCTGGAAAATCAACCGAAGTTTTA
SEQ2403	Aagaagggaagccattacaaatcattaagtgacctctctggaaaatcaaccgaagtttta
SEQ2404	Aagaaggggagccattacaaatcattaagtgacctctctggaaaatcaaccgaagtttta
SEQ2405	AAGAAGGGGAGCCATTACAAATCATTAAGTGACCTCTCTGGAAAATCAACCGAAGTTTTA
SEQ2406	AAGAAGGGGAGCCATTACAAATCATTAAGTGACCTCTCTGGAAAATCAACCGAAGTTTTA
SEQ2407	AAGAAGGGGAGCCATTACAAATCATTAAGTGACCTCTCTGGAAAATCAACAGAAGTTTTA
SEQ2408	AAGAAGGGGAGCCATTACAAATCATTAAGTGACCTCTCTGGAAAATCAACCGAAGTTTTA
-	
SEQ2409	AAGAAGGGGAGCCATTACAAATCATTAAGTGACCTCTCTGGAAAATCAACCGAAGTTTTA
SEQ2410	AAGAAGGGGAGCCATTACAAATCATTAAGTGACCTCTCTGGAAAATCAACAGAAGTTTTA
SEQ2411	AAGAAGGGGAGCCATTACAAATCATTAAGTGACCTCTCTGGAAAATCAACAGAAGTTTTA
SEQ2401	TCTGGCGTTAACTATGCACAGGTTCTAGAAAATTGGAATAAAAATCATCCTAATAAAAA
SEQ2402	TCTGGCGTTAACTATGCACAGGTTCTAGAAAATTGGAATAAAAATCATCCTAATAAAAA
SEQ2403	TCTGGCGTTAACTATGCACAGGTTCTAGAAAATTGGAATAAAAATCATCCTAATAAAAAA
SEQ2404	
	TCTGGCGTTAACTATGCACAGGTTCTAGAAAATTGGAATAAAAATCATCCTAATAAAAA
SEQ2405	TCTGGCGTTAACTATGCACAGGTTCTAGAAAATTGGAATAAAAATCATNNTAATAAAAA
SEQ2406	TCTGGCGTTAACTATGCACAGGTTCTAGAAAATTGGAATAAAAATCATCCTAATAAAAAA
SEQ2407	TCTGGCGTTAACTATGCACAGGTTCTAGAAAATTGGAATAAAAATCATCCTAATAAAAA
SEQ2408	TCTGGCGTTAACTATGCACAGGTTCTAGAAAATTGGAATAAAAATCATCCTAATAAAAA
SEQ2409	TCTGGCGTTAACTATGCACAGGTTCTAGAAAATTGGAATAAAAATCATCCTAATAAAAA
SEQ2410	TCTGGCGTTAACTATGCACAGGTTCTAGAAAATTGGAATAAAAATCATCCTAATAAAAAA
-	
SEQ2411	TCTGGCGTTAACTATGCACAGGTTCTAGAAAATTGGAATAAAAATCATCCTAATAAAAA
SEQ2401	CCAATAAAAATCAAATATGTTTCTGGGACAACTGGTGTTACTAGCAGATTAAAAAATATT
SEQ2402	CCAATAAAATCAAATATGTTTCTGGGACAACTGGTGTTACTAGCAGATTAAAAAATATT
SEQ2403	CCAATAAAAATCAAATATGTTTCTGGGACAACTGGTGTTACTAGCAGATTAAAAAATATT
SEQ2404	CCAATAAAAATCAAATATGTTTCTGGGACAACTGGTGTTACTAGCAGATTAAAAAATATT
_	CONTRACTOR AND AND AND AND CONTRACTOR OF A STATE OF A S
SEQ2405	CCANTAAAAATNAAATATGTTTCTGGGACAACTGGTGTTACTAGCAGATTAAAAAATATT
SEQ2406	CCAATAAAAATCAAATATGTTTCTGGGACAACTGGTGTTACTAGCAGATTAAAAAATATT
SEQ2407	CCAATAAAAATCAAATATGTTTCTGGGACAACTGGTGTTACTAGCAGATTAAAAAATATT
SEQ2408	CCAATAAAAATCAAATATGTTTCTGGGACAACTGGTGTTACTAGCAGATTAAAAAATATT
SEQ2409	CCAATAAAAATCAAATATGTTTCTGGGACAACTGGTGTTACTAGCAGATTAAAAAATATT
_	A



SEQ2410 SEQ2411	CCAATAAAAATCAAATATGTTTCTGGGACAACTGGTGTTACTAGCAGATTAAAAAATATT CCAATAAAAATCAAATATGTTTCTGGGACAACTGGTGTTACTAGCAGATTAAAAAATATT
SEQ2401	GAGAGTGGGAAAATTGACTTTATCCTATATGATGCCATTTCATCTGACTATATTGTAAAA
SEQ2402	GAGAGTGGGAAAATTGACTTTATCCTATATGATGCCATTTCATCCGACTATATTGTAAAA
SEQ2403	GAGAGTGGGAAAATTGACTTTATCCTATATGATGCCATTTCATCCGACTATATTGTAAAA
SEQ2404	GAGAGTGGGAAAATTGACTTTATCCTATATGATGCCATTTCATCCGACTATATTGTAAAA
SEQ2405	GAGAGTGGGAAAATTGACTTTATCCTATATGATGCCATTTCATCCGACTATATTGTAAAA
SEQ2406	GAGAGTGGGAAAATTGACTTTATCCTATATGATGCCATTTCATCCGACTATATTGTAAAA
SEQ2407	GAGAGTGGAAAAATTGACTTTATCCTATATGATGCCATTTCATCTGACTATATTGTAAAA
SEQ2408	GAGAGTGGGAAAATTGACTTTATCCTATATGATGCCATTTCATCCGACTATATTGTAAAA
SEQ2409	GAGAGTGGGAAAATTGACTTTATCCTATATGATGCCATTTCATCCGACTATATTGTAAAA
SEQ2410	GAGAGTGGAAAAATTGACTTTATCCTATATGATGCCATTTCATCTGACTATATTGTAAAA
SEQ2411	GAGAGTGGAAAAATTGACTTTATCCTATATGATGCCATTTCATCTGACTATATTGTAAAA
SEQ2401	GATCAATCATTAAACTTAAGCGTTTCTCCTTTGAAAGGTAAAATTGGTAATAATAAGGAT
SEQ2402	GACCAATCATTAAACTTAAGCGTTTCTCCTTTGAAAGGTAAAATTGGTAATAATAAGGAT
SEQ2403	GACCAATCATTAAACTTAAGCGTTTCTCCTTTGAAAGGTAAAATTGGTAATAATAAGGAT
SEQ2404	GACCAATCATTAAACTTAAGCGTTTCTCCTTTGAAAGGTAAAATTGGTAATAATAAGGAT
SEQ2405	GACCAATCATTAAACTTAAGCGTTTCTCCTTTGAAAGGTAAAATTGGTAATAATAAGGAT
SEQ2406	GACCAATCATTAAACTTAAGCGTTTCTCCTTTGAAAGGTAAAATTGGTAATAATAAGGAT
SEQ2407	GATCAATCATTAAACTTAAGCGTTTCTCCTTTGAAAGGTAAAATTGGTAATAATAAGGAT
SEQ2408	GACCAATCATTAAACTTAAGCGTTTCTCCTTTGAAAGGTAAAATTGGTAATAATAAGGAT
SEQ2409	GACCAATCATTAAACTTAAGCGTTTCTCCTTTGAAAGGTAAAATTGGTAATAATAAGGAT
SEQ2410	GATCAATCATTAAACTTAAGCGTTTCTCCTTTGAAAGGTAAAATTGGTAATAATAAGGAT GATCAATCATTAAACTTAAGCGTTTCTCCTTTGAAAGGTAAAATTGGTAATAATAAGGAT
SEQ2411	
SEQ2401	GGATTAGAATACCTCCTTTTACCAAAAGATAAAAAAGGTAAAACTCTACAGAAATTTATA
SEQ2402	GGACTAGAATACCTCCTTTTACCAAAAGATAAAAAAGGTAAAACTCTACAGAAATTTATA
SEQ2403	GGACTAGAATACCTCCTTTTACCAAAAGATAAAAAAGGTAAAAACTCTACAGAAATTTATA GGACTAGAATACCTCCTTTTACCAAAAGATAAAAAAGGTAAAAACTCTACAGAAATTTATA
SEQ2404 SEQ2405	GGACTAGAATACCTCCTTTTACCAAAAGATAAAAAAGGTAAAACTCTACAGAAATTTATA GGACTAGAATACCTCCTTTTTACCAAAAGATAAAAAAGGTAAAACTCTACAGAAATTTATA
SEQ2405	GGACTAGAATACCTCCTTTTACCAAAAGATAAAAAAGGTAAAACTCTACAGAAATTTATA
SEQ2407	GGATTAGAATACCTCCTTTTACCAAAAGATAAAAAAGGTAAAACTCTACAGAAATTTATA
SEQ2408	GGACTAGAATACCTCCTTTTACCAAAAGATAAAAAAGGTAAAACTCTACAGAAATTTATA
SEQ2409	GGACTAGAATACCTCCTTTTACCAAAAGATAAAAAAGGTAAAACTCTACAGAAATTTATA
SEQ2410	GGATTAGAATACCTCCTTTTACCAAAAGATAAAAAAGGTAAAACTCTACAGAAATTTATA
SEQ2411	GGATTAGAATACCTCCTTTTACCAAAAGATAAAAAAGGTAAAACTCTACAGAAATTTATA
SEQ2401	ATAAGCGTATTAAAGTTTTGAAAGAAGATGGTACTTTGGCACGTTTAAGTAAACAATAT
SEQ2402	ATAAGCGTATTAAAGTTTTGAAAGAAAATGGTACTTTGGCACGTTTAAGTAAACAATAT
SEQ2403	
SEQ2404	ATAAGCGTATTAAAGTTTTGAAAGAAAATGGTACTTTGGCACGTTTAAGTAAACAATAT
SEQ2405	ATAAGCGT
SEQ2406	ATAAGCGTATTAAAGTTTTGAAAGAAAATGGTACTTTGGCACGTTTAAGTAAACAATAT
SEQ2407	ATAAGCGTATTAAAGTTTTGAAAGAAGATGGTACTTTGGCACGTTTAAGTAAACAATAT
SEQ2408	ATAAGCGTATTAAAGTTTTGAAAGAAAATGGTACTTTGGCACGTTTAAGTAAACAATAT
SEQ2409	ATAAGCGTAATAAAGTTTTGAAAGAAAATGGTA
SEQ2410	ATAAGCGTATTAAAGTTTTGAAAGAAGATGGTACTTTGGCACGTTTAAGTAAACAATAT
SEQ2411	ATAAGCGTATTAAAGTTTTGAAAGAAGATGGTACTTTGGCACGTTTAAGTAAACAATAT
SEQ2401	TCGGTGGAGATTACGTTTCAAACATTGATAAA
SEQ2402	TCGGTGGAGATTACGTTTCAAACATTGATAAA
SEQ2403	
SEQ2404	TCGGTGGAGATTACGTTTCAAACATTGATAAA
SEQ2405	
SEQ2406	TCGGTGGAGATTACGTTTCAAACATTGATAAA
SEQ2407	TCGGTGGAGATTACGTTTCAAACATTGATAAA
SEQ2408	TCGGTGGAGATTACGTTTCAAACATTGATAAA
SEQ2409 SEQ2410	TCGGTGGAGATTACGTTTCAAACATTGATAAA
SEQ2410 SEQ2411	TCGGTGGAGATTACGTTTCAAACATTGATAAAGTRCMARATVSTNCSRATNGTSAGABC
ange	199919999111091111091919991199119999119999

## Figure 24: C mparative Sequences relating to SAG0290 (ABC transporter, substrate-binding protein)

SEQ2401	
SEQ2402	
SEQ2403	
SEQ2404	
SEQ2405	
SEQ2406	
SEQ2407	
SEQ2408	
SEQ2409	
SEQ2410	~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~
SEQ2411	RANSRTRSTBSTRATBNDNGRTN

#### >SEQ ID NO 2450: 8\_1169NT frame: 1

VSVQASEKVELKVATDSDTAPFTYQKDGKFKGYDVDVVKAVFKGSKYKVTFKTVPFDTIS TGIDAGKFDLSANDFSYNKERAEKYLFSDPISRSNYAVVGKKGSHYKSLSDLSGKSTEVL SGVNYAQVLENWNKNHPNKKPIKIKYVSGTTGVTSRLKNIESGKIDFILYDAISSDYIVK DQSLNLSVSPLKGKIGNNKDGLEYLLLPKDKKGKTLQKFINKRIKVLKEDGTLARLSKQY FGGDYVSNIDK

#### >SEQ ID NO 2451:8\_18RS21 frame: 1

VSVQASEKVELKVATDSDTAPFTYXKDGKFKGYDVDVVKAVFKGSKYKVTFKTVPFDTIS TGIDAGKFDLSANDFSYNKERAEKYLFSDPISRSNYAVVGKKGSHYKSLSDLSGKSTEVL SGVNYAQVLENWNKNHPNKKPIKIKYVSGTTGVTSRLKNIESGKIDFILYDAISSDYIVK DQSLNLSVSPLKGKIGNNKDGLEYLLLPKDKKGKTLQKFINKRIKVLKENGTLARLSKQY FGGDYVSNIDK

#### >SEQ ID NO 2452:8 2603 frame: 2

FKGYDVDVVKAVFKGSKYKVTFKTVPFDTISTGIDAGKFDLSANDFSYNKERAEKYLFSD PISRSNYAVVGKKGSHYKSLSÓLSGKSTEVLSGVNYAQVLENWNKNHPNKKPIKIKYVSG TTGVTSRLKNIESGKIDFILYDAISSDYIVKDQSLNLSVSPLKGKIGNNKDGLEYLLLPK DKK

#### >SEQ ID NO 2453:8\_090 frame: 1

VSVQASEKVELKVATDSDTAPFTYQKDGKFKGYDVDVVKAVFKGSKYKVTFKTVPFDTIS TGIDAGKFDLSANDFSYNKERAEKYLFSDPISRSNYAVVGKKGSHYKSLSDLSGKSTEVL SGVNYAQVLENWNKNHPNKKPIKIKYVSGTTGVTSRLKNIESGKIDFILYDAISSDYIVK DQSLNLSVSPLKGKIGNNKDGLEYLLLPKDKKGKTLQKFINKRIKVLKENGTLARLSKQY FGGDYVSNIDK

#### >SEQ ID NO 2454:8 A909 frame: 1

VSVQASEKVELKVATDSDTAPFTYQKDGKFKGYDVDVVKAVFKGSKYKVTFKTVPFDTIS TGIDAGKFDLSANDFSYNKERAEKYLFSDPISRSNYAVVGKKGSHYKSLSDLSGKSTEVL SGVNYAQVLENWNKNHXNKKPXKXKYVSGTTGVTSRLKNIESGKIDFILYDAISSDYIVK DQSLNLSVSPLKGKIGNNKDGLEYLLLPKDKKGKTLQKFINKR

#### >SEQ ID NO 2455: 8 CJB110 frame: 1

VSVQASEKVELKVATDSDTAPFTYQKDGKFKGYDVDVVKAVFKGSKYKVTFKTVPFDTIS TGIDAGKFDLSANDFSYNKERAEKYLFSDPISRSNYAVVGKKGSHYKSLSDLSGKSTEVL SGVNYAQVLENWNKNHPNKKPIKIKYVSGTTGVTSRLKNIESGKIDFILYDAISSDYIVK DQSLNLSVSPLKGKIGNNKDGLEYLLLPKDKKGKTLQKFINKRIKVLKENGTLARLSKQY FGGDYVSNIDK

#### >SEQ ID NO 2456: 8 COH1 frame: 1

VSVQASEKVELKVATDSDTAPFTYQKDGKFKGYDVDVVKAVFKGSKYKVTFKTVPFDTIS TGIDAGKFDLSANDFSYNKERAEKYLFSDPISRSNYAVVGKKGSHYKSLSDLSGKSTEVL SGVNYAQVLENWNKNHPNKKPIKIKYVSGTTGVTSRLKNIESGKIDFILYDAISSDYIVK DQSLNLSVSPLKGKIGNNKDGLEYLLLPKDKKGKTLQKFINKRIKVLKEDGTLARLSKQY FGGDYVSNIDK

#### >SEQ ID NO 2457:8\_H36B frame: 1

VSVQASEKVELKVATDSDTAPFTYQKDGKFKGYDVDVVKAVFKGSKYKVTFKTVPFDTIS TGIDAGKFDLSANDFSYNKERAEKYLFSDPISRSNYAVVGKKGSHYKSLSDLSGKSTEVL SGVNYAQVLENWNKNHPNKKPIKIKYVSGTTGVTSRLKNIESGKIDFILYDAISSDYIVK DQSLNLSVSPLKGKIGNNKDGLEYLLLPKDKKGKTLQKFINKRIKVLKENGTLARLSKQY FGGDYVSNIDK



## Figure 24: C mparative Sequences relating t SAG0290 (ABC transporter, substrate-binding protein)

#### >SEQ ID NO 2458:8 JM9130013 frame: 1

VSVQASEKVELKVATDSDTAPFTYQKDGKFKGYDVDVVKAVFKGSKYKVTFKTVPFDTIS TGIDAGKFDLSANDFSYNKERAEKYLFSDPISRSNYAVVGKKGSHYKSLSDLSGKSTEVL SGVNYAQVLENWNKNHPNKKPIKIKYVSGTTGVTSRLKNIESGKIDFILYDAISSDYIVK DQSLNLSVSPLKGKIGNNKDGLEYLLLPKDKKGKTLQKFINKRNKVLKENG

#### >SEQ ID NO 2459:8 M732 frame: 1

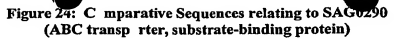
VSVQASEKVELKVATDSDTAPFTYQKDGKFKGYDVDVVKAVFKGSKYKVTFKTVPFDTIS TGIDAGKFDLSANDFSYNKERAEKYLFSDPISRSNYAVVGKKGSHYKSLSDLSGKSTEVL SGVNYAQVLENWNKNHPNKKPIKIKYVSGTTGVTSRLKNIESGKIDFILYDAISSDYIVK DQSLNLSVSPLKGKIGNNKDGLEYLLLPKDKKGKTLQKFINKRIKVLKEDGTLARLSKQY FGGDYVSNIDK

#### >SEQ ID NO 2460:8\_M781 frame: 1

VSVQASEKVELKVATDSDTAPFTYQKDGKFKGYDVDVVKAVFKGSKYKVTFKTVPFDTIS TGIDAGKFDLSANDFSYNKERAEKYLFSDPISRSNYAVVGKKGSHYKSLSDLSGKSTEVL SGVNYAQVLENWNKNHPNKKPIKIKYVSGTTGVTSRLKNIESGKIDFILYDAISSDYIVK DQSLNLSVSPLKGKIGNNKDGLEYLLLPKDKKGKTLQKFINKRIKVLKEDGTLARLSKQY FGGDYVSNIDK

SEQ2450	SVQASEKVELKVATDSDTAPFTYQKDGKFKGYDVDVVKAVFKGSKYKVTFKTVPFDTIS
SEQ2451	SVQASEKVELKVATDSDTAPFTYXKDGKFKGYDVDVVKAVFKGSKYKVTFKTVPFDTIS
SEQ2452	FKGYDVDVVKAVFKGSKYKVTFKTVPFDTIS
SEQ2453	SVQASEKVELKVATDSDTAPFTYQKDGKFKGYDVDVVKAVFKGSKYKVTFKTVPFDTIS
SEQ2454	SVQASEKVELKVATDSDTAPFTYQKDGKFKGYDVDVVKAVFKGSKYKVTFKTVPFDTIS
SEQ2455	SVQASEKVELKVATDSDTAPFTYQKDGKFKGYDVDVVKAVFKGSKYKVTFKTVPFDTIS
SEQ2456	SVQASEKVELKVATDSDTAPFTYQKDGKFKGYDVDVVKAVFKGSKYKVTFKTVPFDTIS
SE02457	SVQASEKVELKVATDSDTAPFTYQKDGKFKGYDVDVVKAVFKGSKYKVTFKTVPFDTIS
SEQ2458	SVQASEKVELKVATDSDTAPFTYQKDGKFKGYDVDVVKAVFKGSKYKVTFKTVPFDTIS
SEQ2459	
SE02460	SVQASEKVELKVATDSDTAPFTYQKDGKFKGYDVDVVKAVFKGSKYKVTFKTVPFDTIS
3EQ2460 .	SVQASEKVELKVATDSDTAPFTYQKDGKFKGYDVDVVKAVFKGSKYKVTFKTVPFDTIS
200450	
SEQ2450	TGIDAGKFDLSANDFSYNKERAEKYLFSDPISRSNYAVVGKKGSHYKSLSDLSGKSTEVL
SEQ2451	TGIDAGKFDLSANDFSYNKERAEKYLFSDPISRSNYAVVGKKGSHYKSLSDLSGKSTEVL
SEQ2452	TGIDAGKFDLSANDFSYNKERAEKYLFSDPISRSNYAVVGKKGSHYKSLSDLSGKSTEVL
SEQ2453	TGIDAGKFDLSANDFSYNKERAEKYLFSDPISRSNYAVVGKKGSHYKSLSDLSGKSTEVL
SEQ2454	TGIDAGKFDLSANDFSYNKERAEKYLFSDPISRSNYAVVGKKGSHYKSLSDLSGKSTEVL
SEQ2455	TGIDAGKFDLSANDFSYNKERAEKYLFSDPISRSNYAVVGKKGSHYKSLSDLSGKSTEVL
SEQ2456	TGIDAGKFDLSANDFSYNKERAEKYLFSDPISRSNYAVVGKKGSHYKSLSDLSGKSTEVL
SEQ2457	TGIDAGKFDLSANDFSYNKERAEKYLFSDPISRSNYAVVGKKGSHYKSLSDLSGKSTEVL
SEQ2458	TGIDAGKFDLSANDFSYNKERAEKYLFSDPISRSNYAVVGKKGSHYKSLSDLSGKSTEVL
SEQ2459	TGIDAGKFDLSANDFSYNKERAEKYLFSDPISRSNYAVVGKKGSHYKSLSDLSGKSTEVL
SEQ2460	TGIDAGKFDLSANDFSYNKERAEKYLFSDPISRSNYAVVGKKGSHYKSLSDLSGKSTEVL
SEQ2450	SGVNYAQVLENWNKNHPNKKPIKIKYVSGTTGVTSRLKNIESGKIDFILYDAISSDYIVK
SE02451	SGVNYAQVLENWNKNHPNKKPIKIKYVSGTTGVTSRLKNIESGKIDFILYDAISSDYIVK
SEQ2452	SGVNYAQVLENWNKNHPNKKPIKIKYVSGTTGVTSRLKNIESGKIDFILYDAISSDYIVK
SEQ2453	SGVNYAQVLENWNKNHPNKKPIKIKYVSGTTGVTSRLKNIESGKIDFILYDAISSDYIVK
SEQ2454	SGVNYAQVLENWNKNHXNKKPXKXKYVSGTTGVTSRLKNIESGKIDFILYDAISSDYIVK
SE02455	SGVNYAQVLENWNKNHPNKKPIKIKYVSGTTGVTSRLKNIESGKIDFILYDAISSDYIVK
SEQ2456	SGVNYAQVLENWNKNHPNKKPIKIKYVSGTTGVTSRLKNIESGKIDFILYDAISSDYIVK
SEQ2457	SGVNYAQVLENWNKNHPNKKPIKIKIVSGIIGVISKLKNIESGKIDFILIDAISSDIIVK
SEQ2457	
SEQ2456 SEQ2459	SGVNYAQVLENWNKNHPNKKPIKIKYVSGTTGVTSRLKNIESGKIDFILYDAISSDYIVK
	SGVNYAQVLENWNKNHPNKKPIKIKYVSGTTGVTSRLKNIESGKIDFILYDAISSDYIVK
SEQ2460	SGVNYAQVLENWNKNHPNKKPIKIKYVSGTTGVTSRLKNIESGKIDFILYDAISSDYIVK
SEQ2450	DQSLNLSVSPLKGKIGNNKDGLEYLLLPKDKKGKTLQKFINKRIKVLKEDGTLARLSKQY
SEQ2451	DQSLNLSVSPLKGKIGNNKDGLEYLLLPKDKKGKTLQKFINKRIKVLKENGTLARLSKQY
SEQ2452	DQSLNLSVSPLKGKIGNNKDGLEYLLLPKDKK
SEQ2453	DQSLNLSVSPLKGKIGNNKDGLEYLLLPKDKKGKTLQKFINKRIKVLKENGTLARLSKQY
SEQ2454	DQSLNLSVSPLKGKIGNNKDGLEYLLLPKDKKGKTLQKFINKR
SEQ2455	DQSLNLSVSPLKGKIGNNKDGLEYLLLPKDKKGKTLQKFINKRIKVLKENGTLARLSKQY
SEQ2456	DQSLNLSVSPLKGKIGNNKDGLEYLLLPKDKKGKTLQKFINKRIKVLKEDGTLARLSKQY
SEQ2457	DQSLNLSVSPLKGKIGNNKDGLEYLLLPKDKKGKTLQKFINKRIKVLKENGTLARLSKQY
SEQ2458	DQSLNLSVSPLKGKIGNNKDGLEYLLLPKDKKGKTLOKFINKRNKVLKENG
SEQ2459	DQSLNLSVSPLKGKIGNNKDGLEYLLLPKDKKGKTLOKFINKRIKVLKEDGTLARLSKOY
SEQ2460	DQSLNLSVSPLKGKIGNNKDGLEYLLLPKDKKGKTLQKFINKRIKVLKEDGTLARLSKQY





SEQ2450	GGDYVSNIDK
SEQ2451	<b>GGDYVSNIDK</b>
SEQ2452	~~~~~~
SEQ2453	GGDYVSNIDK
SEQ2454	
SEQ2455	GGDYVSNIDK
SEQ2456	GGDYVSNIDK
SEQ2457	GGDYVSNIDK
SEQ2458	
SEQ2459	GGDYVSNIDK
SEQ2460	GGDYVSNIDK

## Table 25? Comparative Sequences relating to SAGU508 (protein of unknown function)

SEQ ID NO. 2501: SAG0368 FROM THE 090 GBS TYPE IA STRAIN GAAACAAAGCCGTTTTCAATACTATTAATGGGGGTGGACACAGGTTCAGAGCATCGAAAATCTAAGTGGTCAGGAAAT AGCGATTCTATGATCTTAGTCACTATAAATCCTAAAACTAATAAAACAACGATGACAAGCTTAGAACGTGACGTATTG ATTAAATTGAGTGGTCCCAAAAATAATGGACAGACTGGAGTAGAAGCCAAAGCTAAATGCAGCCTATGCTTCTGGTGGT GCGGAAATGGCATTGATGACTGTTCAAGACTTATTAGATATTAATGTTGATTACTTTATGCAAATTAATATGCAAGGA TTAGTTGATTTAGTCAATGCTGTTGGTGGTATAACAGTAACTAATAAATTTGACTTTCCAATATCAATTGCTGCCAAT CGCTATGATGATCCAGAGGGAGATTATGGGCGTCAAAAAAGGACAACGTGAAGTAATTCAAAAAAGTCCTTAAAAAAATA TCATCAAAAACGATTCCTAATTTGTTAGCTTATAAAGATTCATTGGAACATATTAAATCTTATCAGTTGAAGGGTGAA GACGCTACTTTATCAGATGGTGGCTCTTATCAAATTTTAACTAAGAAACATCTACTTGCAGTTCAAAATAGAATTAAG AAAGAACTGGATAAAAAGCGTAGTAAAACTCTGAAGACAAGCGCGATTCTATATGAAGATTACTATGGTACTACTGCT AGTAATGATTCTTCTACTTATTCATCAACACAGAGAATAATTATAATACAACACCTTATTCAGAAGCACCACCAAGT TACAGTGGTAATACTACTTATAGTTCTGAGACTAATCAAACACTCATCAAAATTACTATAATAGTAGCACTCCTGCT AGTAACTATAGCAGTAACACTAACACAGGTCAGGCTGATTCAAGTGGAAGTGTCAATAATCATAACGGGGCTGCAACG CCTAATCCA

SEQ ID NO. 2502: SAG0368 FROM THE 1169NT1 GBS TYPE V STRAIN GAAACAAAGCCGTTTTCAATACTATTAATGGGGGTGGACACAGGTTCAGAGCATCGAAAATCTAAGTTGGTCAGGAAA TAGCGATTCTATGATCTTAGTCACTATAAATCCTAAAACTAATAAAACAACGATGACAAGCTTAGAACGTGACGTATT GATTAAATTGAGTGGTCCCAAAAATAATGGACAGACTGGCGTAGAAGCCAAAGCTAAATGCAGCCTATGCTTCTGGTGG TGCGGAAATGGCATTGATGACTGTTCAAGACTTATTAGATATTAATGTTGATTACTTTATGCAAATTAATATGCAAGG ATTAGTTGATTTAGTCAATGCTGTTGGTGGTATAACAGTAACTAATAAATTTGACTTTCCAATATCAATTGCTGCCAA GCGCTATGATGATCCAGAGGGAGATTATGGGCGTCAAAAAAGACAACGTGAAGTAATTCAAAAAGTCCTTAAAAAAAT ATCATCAAAAACGATTCCTAATTTGTTAGCTTATAAAGATTCATTGGAACATATTAAATCTTATCAGTTGAAAGGTGA AGACGCTACTTTATCAGATGGTGGCTCTTATCAAATTTTAACTAAGAAACATCTACTTGCAGTTCAAAATAGAATTAA GAAAGAACTAGATAAAAAGCGTAGTAAAACTCTGAAGACAAGCGCGATTCTATATGAAGATTACTATGGTACTACTGC TAGTAATGATTCTTCTACTTATTCATCAACACAAGAGAATAATTATAATACAACACCTTATTCAGAAGCACCAAG TTACAGTGGTAATACTACTTATAGTTCTGAGACTAATCAAACAACTCATCAAAGTTACTATAATAGTAGCACTCCTGC TAATAACTATAGCAGTAACACTAACACAGGTCAGGCTGATTCAAGTGGAAGTGTCAATAATCATAATGGGGCTGCAAC **GCCTAATCCA** 

#### SEQ ID NO. 2503 SAG0368 FROM THE 18RS21 GBS TYPE II STRAIN

GAAACAAAGCCGTTTTCAATACTATTAATGGGGGTGGACACAGGTTCAGAGCATCGAAAATCTAAGTGGTCAGGAAAT AGCGATTCTATGATCTTAGTCACTATAAATCCTAAAACTAATAAAACAACGATGACAAGCTTAGAACGTGACGTATTG ATTAAATTGAGTGGTCCCAAAAATAATGGACAGACTGGAGTAGAAGCAAAGCTAAATGCAGCCTATGCTTCTGGTGGT GCGGAAATGGCATTGATGACTGTTCAAGACTTATTAGATATTAATGTTGATTACTTTATGCAAATTAATATGCAAGGA TTAGTTGATTTAGTCAATGCTGTTGGTGGTATAACAGTAACTAATAAATTTGACTTTCCAATATCAATTGCTGCCAAT CGCTATGATGATCCAGAGGGAGATTATGGGCGTCAAAAAAGACAACGTGAAGTAATTCAAAAAAGTCCTTAAAAAAATA TCATCAAAAACGATTCCTAATTTGTTAGCTTATAAAGATTCATTGGAACATATTAAATCTTATCAGTTGAAGGGTGAA GACGCTACTTTATCAGATGGTGGCTCTTATCAAATTTTAACTAAGAAACATCTACTTGCAGTTCAAAATAGAATTAAG AAAGAACTGGATAAAAAGCGTAGTAAAACTCTGAAGACAAGCGCGATTCTATATGAAGATTACTATGGTACTACTGCT AGTAATGATTCTTCTACTTATTCATCAACACAAGAGAATAATTATAATACAACACCTTATTCAGAAGCACCACCAAGT TACAGTGGTAATACTACTTATAGTTCTGAGACTAATCAAACAACTCATCAAAATTACTATAATAGTAGCACTCCTGCT AGTAACTATAGCAGTAACACTAACACAGGTCAGGCTGATTCAAGTGGAAGTGTCAATAATCATAACGGGGCTGCAACG CCTAATCCA



## Table 25: Comparativ Sequences relating t SAG0308 (pr tein f unknown function)

SEQ ID NO. 2504: SAG0368 FROM THE 2603 V/R GBS TYPE V STRAIN GAAACAAAGCCGTTTTCAATACTATTAATGGGGGTGGACACAGGTTCAGAGCATCGAAAATCTAAGTGGTCAGGAAAT AGCGATTCTATGATCTTAGTCACTATAAATCCTAAAACTAATAAAACAACGATGACAAGCTTAGAACGTGACGTATTG ATTAAATTGAGTGGTCCCAAAAATAATGGACAGACTGGAGTAGAAGCAAAGCTAAATGCAGCCTATGCTTCTGGTGGT GCGGAAATGGCATTGATGACTGTTCAAGACTTATTAGATATTAATGTTGATTACTTTATGCAAATTAATATGCAAGGA TTAGTTGATTTAGTCAATGCTGTTGGTGGTATAACAGTAACTAATAAATTTGACTTTCCAATATCAATTGCTGCCAAT CGCTATGATGATCCAGAGGGAGATTATGGGCGTCAAAAAAGACAACGTGAAGTAATTCAAAAAAGTCCTTAAAAAAATA TCATCAAAAACGATTCCTAATTTGTTAGCTTATAAAGATTCATTGGAACATATTAAATCTTATCAGTTGAAGGGTGAA GACGCTACTTTATCAGATGGTGGCTCTTATCAAATTTTAACTAAGAAACATCTACTTGCAGTTCAAAATAGAATTAAG **AAAGAACTGGATAAAAAGCGTAGTAAAACTCTGAAGACAAGCGCGATTCTATATGAAGATTACTATGGTACTACTGCT** AGTAATGATTCTTCTACTTATTCATCAACACAAGAGAATAATTATAATACAACACCTTATTCAGAAGCACCACCAAGT TACAGTGGTAATACTACTTATAGTTCTGAGACTAATCAAACAACTCATCAAAATTACTATAATAGTAGCACTCCTGCT AGTAACTATAGCAGTAACACTAACACAGGTCAGGCTGATTCAAGTGGAAGTGTCAATAATCATAACGGGGCTGCAACG CCTAATCCA

#### SEQ ID NO. 2505: SAG0368 FROM THE A909 GBS TYPE IA STRAIN GAAACAAAGCCGTTTTCAATACTATTAATGGGGGTGGACACAGGTTCAGAGCATCGAAAATCTAAGTGGTCAGGAAAT AGCGATTCTATGATCTTAGTCACTATAAATCCTAAAACTAATAAAACAACGATGACAAGCTTAGAACGTGACGTATTG ATTAAATTGAGTGGTCCCAAAAATAATGGACAGACTGGAGTAGAAGCCAAAGCTAAATGCAGCCTATGCTTCTGGTGGT GCGGAAATGGCATTGATGACTGTTCAAGACTTATTAGATATTAATGTTGATTACTTTATGCAAATTAATATGCAAGGA TTAGTTGATTTAGTCAATGCTGTTGGTGGTATAACAGTAACTAATAAATTTGACTTTCCAATATCAATTGCTGCCAAT CGCTATGATGATCCAGAGGGAGATTATGGGCGTCAAAAAAGACAACGTGAAGTAATTCAAAAAAGTCCTTAAAAAAATA TCATCAAAAACGATTCCTAATTTGTTAGCTTATAAAGATTCATTGGAACATATTAAATCTTATCAGTTGAAGGGTGAA GACGCTACTTTATCAGATGGTGGCTCTTATCAAATTTTAACTAAGAAACATCTACTTGCAGTTCAAAATAGAATTAAG AAAGAACTGGATAAAAAGCGTAGTAAAACTCTGAAGACAAGCGCGATTCTATATGAAGATTACTATGGTACTACTGCT AGTAATGATTCTTCTACTTATTCATCAACACAGAGAGAATAATTATAATACAACACCTTATTCAGAAGCACCACCAAGT TACAGTGGTAATACTACTTATAGTTCTGAGACTAATCAAACAACTCATCAAAATTACTATAATAGTAGCACTCCTGCT AGTAACTATAGCAGTAACACTAACACAGGTCAGGCTGATTCAAGTGGAAGTGTCAATAATCATAACGGGGCTGCAACG

## SEQ ID NO. 2506: SAG0368 FROM THE CJB110 GBS NONTYPEABLE STRAIN (REVERSE COMPLEMENT)

**CCTAATCCA** 

GAAACAAAGCCGTTTTCAATACTATTAATGGGGGTGGACACAGGTTCAGAGCATCGAAAATCTAAGTGGTCAGGAAAT AGCGATTCTATGATCTTAGTCACTATAAATCCTAAAACTAATAAAACAACGATGACAAGCTTAGAACGTGACGTATTG ATTAAATTGAGTGGTCCCAAAAATAATGGACAGACTGGAGTAGAAGCCAAAGCTAAATGCAGCCTATGCTTCTGGTGGT GCGGAAATGGCATTGATGACTGTTCAAGACTTATTAGATATTAATGTTGATTACTTTATGCAAATTAATATGCAAGGA TTAGTTGATTTAGTCAATGCTGTTGGTGGTATAACAGTAACTAATAAATTTGACTTTCCAATATCAATTGCTGCCAAT TCATCAAAAACGATTCCTAATTTGTTAGCTTATAAAGATTCATTGGAACATATTAAATCTTATCAGTTGAAGGGTGAA GACGCTACTTTATCAGATGGTGGCTCTTATCAAATTTTAACTAAGAAACATCTACTTGCAGTTCAAAATAGAATTAAG AAAGAACTGGATAAAAAGCGTAGTAAAACTCTGAAGACAAGCGCGATTCTATATGAAGATTACTATGGTACTACTGCT AGTAATGATTCTTCTACTTATTCATCAACACAGAGAGAATAATTATAATACAACACCTTATTCAGAAGCACCACCAAGT TACAGTGGTAATACTACTTATTAGTTCTGAGACTAATCAAACAACTCATCAAAATTACTATAATAGTAGCACTCCTGC TAGTAACTATAGCAGTAACACTAACACGGTCAGGCTGATTCAAGTGGAAGTGTCAATAATCATAACGGGGCTGCAAC **GCCTAATCCA** 



## Table 25: C mparative Sequences relating to SAG0568 (protein of unknown function)

### SEQ ID NO. 2507: SAG0368 FROM THE COH1 GBS TYPE III STRAIN (REVERSE COMPLEMENT)

GATTTTAAGCTAGATAAATCAAAAAGTCATGCTATTGAAGAAACAAAGCCGTTTTCAATACTATTAATGGGTGTGGAC ACAGGTTCAGAGCATCGAAAATCTAAGTGGTCAGGAAATAGCGATTCTATGATCTTAGTCACTATAAATCCTAAAACT AATAAAACAACGATGACAAGCTTAGAACGTGACGTATTGATTAAATTGAGTGGTCCCAAAAATAATGGACAGACTGGC GTAGAAGCAAAGCTAAATGCAGCCTATGCTTCTGGTGGTGCGGAAATGGCATTGATGACTGTTCAAGACTTATTAGAT ATTAATGTTGATTACTTTATGCAAATTAATATGCAAGGATTAGTTGATTTGGTCAATGCTGTTGGTGGTATAACAGTA ACTAATAAATTTGACTTTCCAATATCAATTGCTGCCAATGAACCAGAGTACAAGGCTGTTGTTGAACCAGGGACACAT AAAATAAATGGAGAACAAGCACTTGTTTATTCTCGTATGCGCTATGATGATCCAGAGGGAGATTATGGGCGTCAAAAA AGACAACGTGAAGTAATTCAAAAAAGTCCTTAAAAAAATATTGGCGTTAAATAGTATTAGTTCATACAAAAAAATTCTT TCCGCAGTAAGTAATAACATGCAAACTAATATTGAGATATCATCAAAAACGATTCCTAATTTGTTAGCTTATAAAGAT TCATTGGAACATATTAAATCTTATCAGTTGAAGGGTGAAGACGCTACTCTATCAGATGGTGGCTCTTATCAAATTTTA AGCGCGATTCTATATGAAGATTACTATGGTACTACTGCTAGTAATGATTCTTCTACTTATTCATCAACACAAGAGAAT TATTATTÀTACAACACCCTTATTCAGAAGCACCACCAAGTTACAGTGGTAATACTACTTATAGTTCTGAGACTAATCA AACAACTCATCAAAGTTACTATAATAGTAGCACTCCTGCTAGTAACTATAGCAGTAACACTAACACAGGTCAGGCTGA TTCAAGTGGAAGTGTTAATAATTATAACGGGGCTGCAACGCCTAATCCAAACACAGGAACGCAACCAGTACCAGGTCA **AACTAATCCA** 

#### SEQ ID NO. 2508: SAG0368 FROM THE H36b GBS TYPE Ib STRAIN

#### SEQ ID NO. 2509: SAG0368 FROM THE ?????

### SEQ ID NO. 2510: SAG0368 FROM THE JM9130013 GBS TYPE VIII STRAIN (REVERSE COMPLEMENT)

GAAACAAAGCCGTTTTCAATACTATTAATGGGGGTGGACACAGGTTCAGAGCATCGAAAATCTAAGTGGTCAGGAAAT AGCGATTCTATGATCTTAGTCACTATAAATCCTAAAACTAATAAAACAACGATGACAAGCTTAGAACGTGACGTATTG ATTAAATTGAGTGGTCCCAAAAATAATGGACAGACTGGAGTAGAAGCCAAAGCTAAATGCAGCCTATGCTTCTGGTGGT GCGGAAATGGCATTGATGACTGTTCAAGACTTATTAGATATTAATGTTGATTACTTTATGCAAATTAATATGCAAGGA TTAGTTGATTTAGTCAATGCTGTTGGTGGTATAACAGTAACTAATAAATTTGACTTTCCAATATCAATTTGCTGCCAAT CGCTATGATCACAGAGGGAGATTATGGGCGTCAAAAAAGACAACGTGAAGTAATTCAAAAAAGTCCTTAAAAAAATA TCATCAAAAACGATTCCTAATTTGTTAGCTTATAAAGATTCATTGGAACATATTAAATCTTATCAGTTGAAGGGTGAA GACGCTACTTTATCAGATGGTGGCTCTTATCAAATTTTAACTAAGAAACATCTACTTGCAGTTCAAAATAGAATTAAG AAAGAACTGGATAAAAAGCGTAGTAAAAACTCTGAAGACAAGCGCGATTCTATATGAAGATTACTATGGTACTACTGCT AGTAATGATTCTTCTACTTATTCATCAACACAGAGAATAATTATAATACAACACCTTATTCAGAAGCACCACCAAGT TACAGTGGTAATACTACTTATAGTTCTGAGACTAATCAAACAACTCATCAAAATTACTATAATAGTAGCACTCCTGCT AGTAACTATAGCAGTAACACTAACACAGGTCAGGCTGATTCAAGTGGAAGTGTCAATAATCATAACGGGGCTGCAACG CCTAATCCA



## Table 25: Comparative Sequences relating t SAG0568 (protein of unknown functi n)

### SEQ ID NO. 2511: SAG0368 FROM THE M781 GBS TYPE III STRAIN (REVERSE COMPLEMENT)

SEQ2501	772000007775777777777777777777777777777
SEQ2502	
SEQ2503	
SEQ2504	
SEQ2505	
SEQ2506	
SEQ2507	ATTTTAAGCTAGATAAATCAAAAAGTCATGCTATTGAAGAAACAAAGCCGTTTTCAATA
SEQ2508	
SEQ2509	
SEQ2510	
SEQ2511	TTCAATA
SEQ2501	
SEQ2502	
SEQ2503	
SEQ2504	
SEQ2505	
SEQ2506	
SEQ2507	TATTAATGGGTGTGGACACAGGTTCAGAGCATCGAAAATCTAAGTGGTCAGGAAATAGC
SEQ2508	
SEQ2509	
SEQ2510	
SEQ2511	TATTAATGGGTGTGGACACAGGTTCAGAGCATCGAAAATCTAAGTGGTCAGGAAATAGC
SEQ2501	
SEQ2502	
SEQ2503	
SEQ2504	
SEQ2505	
SEQ2506	
SEQ2507	ATTCTATGATCTTAGTCACTATAAATCCTAAAACTAATAAAACAACGATGACAAGCTTA
SEQ2508	~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~
SEQ2509	
SEQ2510	~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~
SEQ2511	ATTCTATGATCTTAGTCACTATAAATCCTAAAACTAATAAAACAACGATGACAAGCTTA



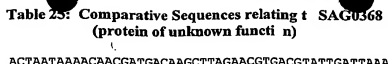
# Table 23: C mparative Sequences relating to SAG0368 (prot in funknown function)

SEQ2501	
SEQ2502	
SEQ2503	
SEQ2504	
SEQ2505	
SEQ2506	
SEQ2507	AACGTGACGTATTGATTAAATTGAGTGGTCCCAAAAATAATGGACAGACTGGCGTAGAA
SEQ2508	
SEQ2509	
SEQ2510	
SEQ2511	AACGTGACGTATTGATTAAATTGAGTGGTCCCAAAAATAATGGACAGACTGGCGTAGAA
SEQ2501	
SEQ2502	
SEQ2503	**
SEQ2504	
SEQ2505	
SEQ2506	
SEQ2507	CAAAGCTAAATGCAGCCTATGCTTCTGGTGGTGCGGAAATGGCATTGATGACTGTTCAA
SEQ2508	
SEQ2509	
SEQ2510	
SEQ2511	CAAAGCTAAATGCAGCCTATGCTTCTGGTGGTGCGGAAATGGCATTGATGACTGȚTCAA
SECOEO1	***************************************
SEQ2501	
SEQ2502	
SEQ2503	
SEQ2504	
SEQ2505	
SEQ2506	
SEQ2507	ACTTATTAGATATTAATGTTGATTACTTTATGCAAATTAATATGCAAGGATTAGTTGAT
SEQ2508	
SEQ2509	
SEQ2510	
SEQ2511	
SEQ2511	ACTTATTAGATATTAATGTTGATTACTTTATGCAAATTAATATGCAAGGATTAGTTGAT
SEQ2501	
SEQ2502	
SEQ2503	
SEQ2504	
SEQ2505	
SEQ2506	
SEQ2507	
	TGGTCAATGCTGTTGGTGGTATAACAGTAACTAATAAATTTGACTTTCCAATATCAATT
SEQ2508	
SEQ2509	
SEQ2510	
SEQ2511	TGGTCAATGCTGTTGGTGGTATAACAGTAACTAATAAATTTGACTTTCCAATATCAATT
SEQ2501	
SEQ2502	
SEQ2502 SEQ2503	
SEQ2504	
SEQ2505	# 0 4 5 5 4 5 5 6 6 6 6 6 6 6 6 6 6 6 6 6 6
SEQ2506	
SEQ2507	CTGCCAATGAACCAGAGTACAAGGCTGTTGTTGAACCAGGGACACATAAAATAAAT
SEQ2508	
SEQ2509	
SEQ2510	, ************************************
SEQ2511	CTGCCAATGAACCAGAGTACAAGGCTGTTGTTGAACCAGGGACACATAAAATAAAT
X	OLOCORIORIONO INCAMOLETE TETTE AACCAGGGACACATAAATAAATGGA



# Table 25: Comparative Sequences relating to SAG0368 (protein of unknown function)

SEQ2501	
SEQ2502	
SEQ2503	
SEQ2504	
SEQ2505	
SEQ2506	
SEQ2507	AACAAGCACTTGTTTATTCTCGTATGCGCTATGATGATCCAGAGGGAGATTATGGGCGT
SEQ2508	
SEQ2509	***************************************
SEQ2510	
SEQ2510	AACAAGCACTTGTTTATTCTCGTATGCGCTATGATGATCCAGAGGGAGATTATGGGCGT
SHQZJII	AACAAGCACIIGIIIAIICICGIAIGCGCIAIGAICAAGGGGAGAIIAIGGGCGT
SE02501	TATAATTTTTCG
SEQ2501	TATATTTTCG
SEQ2502	TATATTTTCG
SEQ2503	TATAATTTTTCG
SEQ2505	TATAATTTTTCG
	TATAATTTTTCG
SEQ2506	TATAATTTTCG
SEQ2507	AAAAAAGACAACGTGAAGTAATTCAAAAAGTCCTTAAAAAAATATTGGCGTTAAATAGT
SEQ2508	TATAATTTTCG
SEQ2509	
SEQ2510	TATAATTTTTCG
SEQ2511	AAAAAAGACAACGTGAAGTAATTCAAAAAGTCCTTAAAAAAATATTGGCGTTAAATAGT
SEQ2501	CTAATGAATTGTCTAAGACTTTTAAAGATTTTAAGCTAGCT
SEQ2502	CTAATGAATTGTCTAAGACTTTTAAAGATTTTAAGCTAGCT
SEQ2503	CTAATGAATTGTCTAAGACTTTTAAAGATTTTAAGCTAGCT
SEQ2504	CTAATGAATTGTCTAAGACTTTTAAAGATTTTAAGCTAGCT
SEQ2505	CTAATGAATTGTCTAAGACTTTTAAAGATTTTAAGCTAGCT
SEQ2506	CTAATGAATTGTCTAAGACTTTTAAAGATTTTAAGCTAGCT
SEQ2507	TTAGTTCAT-ACAAAAAATTCTTTCCGCAGTAAGTAATAACATGCAAACTAATATT
SEQ2508	CTAATGAATTGTCTAAGACTTTTAAAGATTTTAAGCTAGCT
SEQ2509	TTAGTTCAT-ACAAAAAATTCTTTCCGCAGTAAGTAATAACATGCAAACTAATATT
SEQ2510	CTAATGAATTGTCTAAGACTTTTAAAGATTTTAAGCTAGCT
SEQ2511	TTAGTTCAT-ACAAAAAATTCTTTCCGCAGTAAGTAATAACATGCAAACTAATATT
	•
SEQ2501	TTGAAGAAACAAAGCCGTTTTCAATACTATTAATGGGGGTGGACACAGGTTCAGAGCAT
SEQ2502	TTGAAGAAACAAAGCCGTTTTCAATACTATTAATGGGGGTGGACACAGGTTCAGAGCAT
SEQ2503	TTGAAGAAACAAAGCCGTTTTCAATACTATTAATGGGGGTGGACACAGGTTCAGAGCAT
SEQ2504	TTGAAGAAACAAAGCCGTTTTCAATACTATTAATGGGGGTGGACACAGGTTCAGAGCAT
SEQ2505	TTGAAGAAACAAAGCCGTTTTCAATACTATTAATGGGGGTGGACACAGGTTCAGAGCAT
SEQ2506	TTGAAGAAACAAAGCCGTTTTCAATACTATTAATGGGGGTGGACACAGGTTCAGAGCAT
SEQ2507	AGATATCATCAAAAACGATTCCTAATTTGTTAGCTTATAAAGATTCATTGGAACAT
SEQ2508	TTGAAGAAACAAAGCCGTTTTCAATACTATTAATGGGGGTGGACACAGGTTCAGAGCAT
SEQ2509	AGATATCATCAAAAACGATTCCTAATTTGTTAGCTTATAAAGATTCATTGGAACAT
SEQ2510	TTGAAGAAACAAAGCCGTTTTCAATACTATTAATGGGGGTGGACACAGGTTCAGAGCAT
SEQ2511	AGATATCATCAAAAACGATTCCTAATTTGTTAGCTTATAAAGATTCATTGGAACAT
_	
SEQ2501	GAAAATCTAAGT-GGTCAGGAAATAGCGATTCTATGATCTTAGTCACTATAAATCCTAA
SEQ2502	GAAAATCTAAGTTGGTCAGGAAATAGCGATTCTATGATCTTAGTCACTATAAATCCTAA
SEQ2503	GAAAATCTAAGT-GGTCAGGAAATAGCGATTCTATGATCTTAGTCACTATAAATCCTAA
SEQ2504	GAAAATCTAAGT-GGTCAGGAAATAGCGATTCTATGATCTTAGTCACTATAAATCCTAA
SEQ2505	GAAAATCTAAGT-GGTCAGGAAATAGCGATTCTATGATCTTAGTCACTATAAATCCTAA
SEQ2506	GAAAATCTAAGT-GGTCAGGAAATAGCGATTCTATGATCTTAGTCACTATAAATCCTAA
SEQ2507	TTAAATCTTATC-AGTTGAAGGGTGAAGACGCTACTCTATCAG-ATGGTGGCTCTTAT
SEQ2507 SEQ2508	
SEQ2509	GAAAATCTAAGT-GGTCAGGAAATAGCGATTCTATGATCTTAGTCACTATAAATCCTAA
SEQ2510	TTAAATCTTATC-AGTTGAAGGGTGAAGACGCTACTTTATCAG-ATGGTGGCTCTTAT
	GAAAATCTAAGT-GGTCAGGAAATAGCGATTCTATGATCTTAGTCACTATAAATCCTAA
SEQ2511	TTAAATCTTATC-AGTTGAAGGGTGAAGACGCTACTCTATCAGATGGTGGCTCTTAT
SEQ2501	ACTAATAAAACAACGATGACAAGCTTAGAACGTGACGTATTGATTAAATTGAGTGGTCC



	100110111101101001001001001001001001001
SEQ2502	ACTAATAAAACAACGATGACAAGCTTAGAACGTGACGTATTGATTAAATTGAGTGGTCC
SEQ2503	ACTAATAAAACAACGATGACAAGCTTAGAACGTGACGTATTGATTAAATTGAGTGGTCC
SEQ2504	ACTAATAAAACAACGATGACAAGCTTAGAACGTGACGTATTGATTAAATTGAGTGGTCC
SEQ2505	ACTAATAAAACAACGATGACAAGCTTAGAACGTGACGTATTGATTAAATTGAGTGGTCC
SEQ2506	ACTAATAAAACAACGATGACAAGCTTAGAACGTGACGTATTGATTAAATTGAGTGGTCC
SEQ2507	AAATTTTAACTAAGAAACATCTACTTGCAGTTCAAAATAGAATTAAGAAAGA
SEQ2508	ACTAATAAAACAACGATGACAAGCTTAGAACGTGACGTATTGATTAAATTGAGTGGTCC
SEQ2509	AAATTTTAACTAAGAAACATCTACTTGCAGTTCAAAATAGAATTAAGAAAGA
SEQ2510	ACTAATAAAACAACGATGACAAGCTTAGAACGTGACGTATTGATTAAATTGAGTGGTCC
SEQ2511	AAATTTTAACTAAGAAACATCTACTTGCAGTTCAAAATAGAATTAAGAAAGA
SEQ2501	AAAAATAATGGACAGACTGGAGTAGAAGCAAAGCTAAATGCAGCCTATGCTTCTGGT
SEQ2502	AAAAATAATGGACAGACTGGCGTAGAAGCAAAGCTAAATGCAGCCTATGCTTCTGGT
SEQ2503	AAAAATAATGGACAGACTGGAGTAGAAGCAAAGCTAAATGCAGCCTATGCTTCTGGT
SEQ2504	AAAAATAATGGACAGACTGGAGTAGAAGCAAAGCTAAATGCAGCCTATGCTTCTGGT
SEQ2505	AAAAATAATGGACAGACTGGAGTAGAAGCAAAGCTAAATGCAGCCTATGCTTCTGGT
SEQ2506	AAAAATAATGGACAGACTGGAGTAGAAGCAAAGCTAAATGCAGCCTATGCTTCTGGT
SEQ2507	AAAAGCGTAGTAAAACTCTGAAGACAAGCGCGATTCTATATGAAGATTACTATGGTACT
	AAAAATAATGGACAGACTGGAGTAGAAGCAAAGCTAAATGCAGCCTATGCTTCTGGT
SEQ2508	
SEQ2509	AAAAGCGTAGTAAAACTCTGAAGACAAGCGCGATTCTATATGAAGATTACTATGGTACT
SEQ2510	AAAAATAATGGACAGACTGGAGTAGAAGCAAAGCTAAATGCAGCCTATGCTTCTGGT
SEQ2511	AAAAGCGTAGTAAAACTCTGAAGACAAGCGCGATTCTATATGAAGATTACTATGGTACT
SEQ2501	GTGC-GGAAATGGCATTGATGACTGTTCAAGACTTATTAGATATTAATGTTGATTACTT
SEQ2502	GTGC-GGAAATGGCATTGATGACTGTTCAAGACTTATTAGATATTAATGTTGATTACTT
SEQ2503	GTGC-GGAAATGGCATTGATGACTGTTCAAGACTTATTAGATATTAATGTTGATTACTT
SEQ2504	GTGC-GGAAATGGCATTGATGACTGTTCAAGACTTATTAGATATTAATGTTGATTACTT
SEQ2505	GTGC-GGAAATGGCATTGATGACTGTTCAAGACTTATTAGATATTAATGTTGATTACTT
SEQ2506	GTGC-GGAAATGGCATTGATGACTGTTCAAGACTTATTAGATATTAATGTTGATTACTT
SEQ2507	CTGCTAGTAATGATTCTTCTACTTATTCATCAAC-ACAAGAGAATTATTATTAT-ACAA
SEQ2508	GTGC-GGAAATGGCATTGATGACTGTTCAAGACTTATTAGATATTAATGTTGATTACTT
SEQ2509	CTGCTAGTAATGATTCTTCTACTTATTCATCAAC-ACAAGAGAATAATTATAAT-ACAA
SEQ2510	GTGC-GGAAATGGCATTGATGACTGTTCAAGACTTATTAGATATTAATGTTGATTACTT
SEQ2511	CTGCTAGTAATGATTCTTCTACTTATTCATCAAC-ACAAGAGAATAATTATAAT-ACAA
SEQ2501	ATGCAAATTAATATGCAAGGATTAGTTGATTTAGTCAATGCTGTTGGTGGTATAACAGT
SEQ2502	ATGCAAATTAATATGCAAGGATTAGTTGATTTAGTCAATGCTGTTGGTGGTATAACAGT
SEQ2503	ATGCAAATTAATATGCAAGGATTAGTTGATTTAGTCAATGCTGTTGGTGGTATAACAGT
SEQ2504	ATGCAAATTAATATGCAAGGATTAGTTGATTTAGTCAATGCTGTTGGTGGTATAACAGT
SEQ2505	ATGCAAATTAATATGCAAGGATTAGTTGATTTAGTCAATGCTGTTGGTGGTATAACAGT
SEQ2506	ATGCAAATTAATATGCAAGGATTAGTTGATTTAGTCAATGCTGTTGGTGGTATAACAGT
SEQ2507	ACCCTTATTCAGAAGCACCACCAAGTTACAGTGGT-AATACTACTTATAGTTCTGA
SEQ2508	
	ATGCAAATTAATATGCAAGGATTAGTTGATTTAGTCAATGCTGTTGGTGGTATAACAGT
SEQ2509	ACC-TTATTCAGAAGCACCACCAAGTTACAGTGGT-AATACTACTTATAGTTCTGA
SEQ2510	ATGCAAATTAATATGCAAGGATTAGTTGATTTAGTCAATGCTGTTGGTGGTATAACAGT
SEQ2511	ACC-TTATTCAGAAGCACCACCAAGTTACAGTGGT-AATACTACTTATAGTTCTGA
SEQ2501	ACTAATAAATTTGACTTTCCAATATCAATTGCTGCCAATGAACCAGAGTACAAGGCTGT
SEQ2502	ACTAATAAATTTGACTTTCCAATATCAATTGCTGCCAATGAACCAGAGTACAAGGCTGT
SEQ2503	ACTAATAAATTTGACTTTCCAATATCAATTGCTGCCAATGAACCAGAGTACAAGGCTGT
SEQ2504	ACTAATAAATTTGACTTTCCAATATCAATTGCTGCCAATGAACCAGAGTACAAGGCTGT
SEQ2505	ACTAATAAATTTGACTTTCCAATATCAATTGCTGCCAATGAACCAGAGTACAAGGCTGT
SEQ2506	ACTAATAAATTTGACTTTCCAATATCAATTGCTGCCAATGAACCAGAGTACAAGGCTGT
SEQ2507	ACTAATCAAAC-AACTCATCAAAGTTACTAT-AATAGTAGCACTCCTGCTAGT
SEQ2507 SEQ2508	
<del>-</del>	ACTAATAAATTTGACTTTCCAATATCAATTGCTGCCAATGAACCAGAGTACAAGGCTGT
SEQ2509	ACTAATCAAAC-AACTCATCAAAATTACTAT-AATAGTAGCACTCCTGCTAGT
SEQ2510	ACTAATAAATTTGACTTTCCAATATCAATTGCTGCCAATGAACCAGAGTACAAGGCTGT
SEQ2511	ACTAATCAAAC-AACTCATCAAAGTTACTAT-AATAGTAGCACTCCTGCTAGT



# Table 25: Comparative Sequences relating to SAG0368 (protein of unknown function)

SEQ2501	GTTGAACCAGGGACACATAAAATAAATGGAGAACAAGCACTTGTTTATTCTCGTATGCG
SEQ2502	GTTGAACCAGGGACACATAAAATAAATGGAGAACAAGCACTTGTTTATTCTCGTATGCG
SEQ2503	GTTGAACCAGGGACACATAAAATAAATGGAGAACAAGCACTTGTTTATTCTCGTATGCG
SEQ2504	GTTGAACCAGGGACACATAAAATAAATGGAGAACAAGCACTTGTTTATTCTCGTATGCG
SEQ2505	GTTGAACCAGGGACACATAAAATAAATGGAGAACAAGCACTTGTTTATTCTCGTATGCG
SEQ2506	GTTGAACCAGGGACACATAAAATAAATGGAGAACAAGCACTTGTTTATTCTCGTATGCG
SEQ2507	ACTATAGCAGTAACAC-TAACACAGGTCAGGCTGATTCAAGTGGAAGTGTTAATAATTA
SEQ2508	GTTGAACCAGGGACACATAAAATAAATGGAGAACAAGCACTTGTTTATTCTCGTATGCG
SEQ2509	ACTATAGCAGTAACAC-TAACACAGGTCAGGCTGATTCAAGTGGAAGTGTCAATAATCA
SEQ2510	GTTGAACCAGGGACACATAAAATAAATGGAGAACAAGCACTTGTTTATTCTCGTATGCG
SEQ2511	ACTATAGCAGTAACAC-TAACACAGGTCAGGCTGATTCAAGTGGAAGTGTTAATAATTA
224711	NOTHING THE PROPERTY OF THE PR
SEQ2501	TATGATGATCCAGAGGGAGATTATGGGCGTCAAAAAAGACAACGTGAAGTAATTCAAAA
SEQ2502	TATGATGATCCAGAGGGAGATTATGGGCGTCAAAAAAGACAACGTGAAGTAATTCAAAA
SEQ2503	TATGATGATCCAGAGGGAGATTATGGGCGTCAAAAAAGACAACGTGAAGTAATTCAAAA
SEQ2504	TATGATGATCCAGAGGGAGATTATGGGCGTCAAAAAAGACAACGTGAAGTAATTCAAAA
SEQ2505	TATGATGATCCAGAGGGAGATTATGGGCGTCAAAAAAGACAACGTGAAGTAATTCAAAA
SEQ2506	TATGATGATCCAGAGGGAGATTATGGGCGTCAAAAAAGACACGTGAAGTAATTCAAAA
SEQ2507	AACGGGGCTGCAACGCCTAATCCAAACACAGGAACGCAACCAGTACCAGGTCAAACTAA
SEQ2508	TATGATGATCCAGAGGGAGATTATGGGCGTCAAAAAAGACAACGTGAAGTAATTCAAAA
SEQ2509	AACGGGGCTGCAACGCCTAATCCA
SEQ2510	TATGATGATCCAGAGGGAGATTATGGGCGTCAAAAAAGACAACGTGAAGTAATTCAAAA
SEQ2511	AACGGGGCTGCAACGCCTAATCCAAACACAGGAACGCAACCAGTACCAGGTCAAACTAA
	•
SEQ2501	GTCCTTAAAAAAATATTGGCGTTAAATAGTATTAGTTCATACAAAAAAATTCTTTCCGC
SEQ2502	GTCCTTAAAAAAATATTGGCGTTAAATAGTATTAGTTCATACAAAAAAATTCTTTCCGC
SEQ2503	GTCCTTAAAAAAATATTGGCGTTAAATAGTATTAGTTCATACAAAAAAATTCTTTCCGC
SEQ2504	GTCCTTAAAAAAATATTGGCGTTAAATAGTATTAGTTCATACAAAAAAATTCTTTCCGC
SEQ2505	GTCCTTAAAAAAATATTGGCGTTAAATAGTATTAGTTCATACAAAAAAATTCTTTCCGC
SEQ2506	GTCCTTAAAAAAATATTGGCGTTAAATAGTATTAGTTCATACAAAAAAATTCTTTCCGC
SEQ2507	CCA
SEQ2508	GTCCTTAAAAAAATATTGGCGTTAAATAGTA
SEQ2509	
SEQ2510	GTCCTTAAAAAAATATTGGCGTTAAATAGTATTAGTTCATACAAAAAAATTCTTTCCGC
SEQ2511	CCA
SEQ2501	GTAAGTAATAACATGCAAACTAATATTGAGATATCATCAAAAACGATTCCTAATTTGTT
SEQ2501	GTAAGTAATAACATGCAAACTAATATTGAGATATCATCAAAAACGATTCCTAATTTGTT
SEQ2502 SEQ2503	GTAAGTAATAACATGCAAACTAATATTGAGATATCATCAAAAACGATTCCTAATTTGTT
SEQ2504	GTAAGTAATAACATGCAAACTAATATTGAGATATCATCAAAAAACGATTCCTAATTTGTT
SEQ2505	GTAAGTAATAACATGCAAACTAATATTGAGATATCATCAAAAAACGATTCCTAATTTGTT
SEQ2506	
SEQ2506 SEQ2507	GTAAGTAATAACATGCAAACTAATATTGAGATATCATCAAAAACGATTCCTAATTTGTT
SEQ2507	
SEQ2509	
SEQ2509 SEO2510	GTAAGTAATAACATGCAAACTAATATTGAGATATCATCAAAAAACGATTCCTAATTTGTT
SEQ2511	ormorminoni ochmormini i ingnini chi chammooni i colimili di i
SEQ2501	GCTTATAAAGATTCATTGGAACATATTAAATCTTATCAGTTGAAGGGTGAAGACGCTAC
SEQ2502	GCTTATAAAGATTCATTGGAACATATTAAATCTTATCAGTTGAAAGGTGAAGACGCTAC
SEQ2503	GCTTATAAAGATTCATTGGAACATATTAAATCTTATCAGTTGAAGGGTGAAGACGCTAC
SEQ2504	GCTTATAAAGATTCATTGGAACATATTAAATCTTATCAGTTGAAGGGTGAAGACGCTAC
SEQ2505	GCTTATAAAGATTCATTGGAACATATTAAATCTTATCAGTTGAAGGGTGAAGACGCTAC
SEQ2506	GCTTATAAAGATTCATTGGAACATATTAAATCTTATCAGTTGAAGGGTGAAGACGCTAC
SEQ2507	*
SEQ2508	
SEQ2509	*
SEQ2510	GCTTATAAAGATTCATTGGAACATATTAAATCTTATCAGTTGAAGGGTGAAGACGCTAC
SEQ2511	



# Table 25: C mparative Sequences relating to SAG0368 (protein funknown functi n)

SEQ2501	TTATCAGATGGTGGCTCTTATCAAATTTTAACTAAGAAACATCTACTTGCAGTTCAAAA
SEQ2502	TTATCAGATGGTGGCTCTTATCAAATTTTAACTAAGAAACATCTACTTGCAGTTCAAAA
SEQ2503	TTATCAGATGGTGGCTCTTATCAAATTTTAACTAAGAAACATCTACTTGCAGTTCAAAA
SEQ2504	TTATCAGATGGTGGCTCTTATCAAATTTTAACTAAGAAACATCTACTTGCAGTTCAAAA
SEQ2505	TTATCAGATGGTGGCTCTTATCAAATTTTAACTAAGAAACATCTACTTGCAGTTCAAAA
SEQ2506	TTATCAGATGGTGGCTCTTATCAAATTTTAACTAAGAAACATCTACTTGCAGTTCAAAA
SEQ2507	
SEQ2508	
SEQ2509	
SEQ2510	TTATCAGATGGTGGCTCTTATCAAATTTTAACTAAGAAACATCTACTTGCAGTTCAAAA
SEQ2511	
226777	
SEQ2501	AGAATTAAGAAAGAACTGGATAAAAAGCGTAGTAAAACTCTGAAGACAAGCGCGATTCT
SEQ2502	AGAATTAAGAAAGAACTAGATAAAAAGCGTAGTAAAACTCTGAAGACAAGCGCGATTCT
SEQ2502 SEQ2503	AGAATTAAGAAAGAACTGGATAAAAAGCGTAGTAAAACTCTGAAGACAAGCGCGATTCT
SEQ2503 SEQ2504	AGAATTAAGAAAGAACTGGATAAAAAGCGTAGTAAAACTCTGAAGACAAGCGCGATTCT AGAATTAAGAAAAGACTGGATAAAAAGCGTAGTAAAACTCTGAAGACAAGCGCGATTCT
SEQ2505	AGAATTAAGAAAGAACTGGATAAAAAGCGTAGTAAAACTCTGAAGACAAGCGCGATTCT
SEQ2506	AGAATTAAGAAAGAACTGGATAAAAAGCGTAGTAAAACTCTGAAGACAAGCGCGATTCT
SEQ2507	
SEQ2508	
SEQ2509	
SEQ2510	AGAATTAAGAAAGAACTGGATAAAAAGCGTAGTAAAACTCTGAAGACAAGCGCGATTCT
SEQ2511	
SEQ2501	TATGAAGATTACTATGGTACTACTGCTAGTAATGATTCTTCTACTTATTCATCAACACA
SEQ2502	TATGAAGATTACTATGGTACTACTGCTAGTAATGATTCTTCTACTTATTCATCAACACA
SEQ2503	TATGAAGATTACTATGGTACTACTGCTAGTAATGATTCTTCTACTTATTCATCAACACA
SEQ2504	TATGAAGATTACTATGGTACTACTGCTAGTAATGATTCTTCTACTTATTCATCAACACA
SEQ2505	TATGAAGATTACTATGGTACTACTGCTAGTAATGATTCTTCTACTTATTCATCAACACA
SEQ2506	TATGAAGATTACTATGGTACTACTGCTAGTAATGATTCTTCTACTTATTCATCAACACA
SEQ2507	
SEQ2508	
SEQ2509	
SEQ2510	TATGAAGATTACTATGGTACTACTGCTAGTAATGATTCTTCTACTTATTCATCAACACA
SEQ2511	
0085011	
SEQ2501	GAGAATAATTATAATACAACACCTTATTCAGAAGCACCACCAAGTTACAGTGGTAATAC
SEQ2502	GAGAATAATTATAATACAACACCTTATTCAGAAGCACCACCAAGTTACAGTGGTAATAC
SEQ2502	GAGAATAATTATAATACAACACCTTATTCAGAAGCACCACCAAGTTACAGTGGTAATAC
	GAGAATAATTATAATACAACACCTTATTCAGAAGCACCACCAAGTTACAGTGGTAATAC
SEQ2504	
SEQ2505	GAGAATAATTATAATACAACACCTTATTCAGAAGCACCACCAAGTTACAGTGGTAATAC
SEQ2506	GAGAATAATTATAATACAACACCTTATTCAGAAGCACCACCAAGTTACAGTGGTAATAC
SEQ2507	
SEQ2508	
SEQ2509	
SEQ2510	GAGAATAATTATAATACAACACCTTATTCAGAAGCACCAACCA
SEQ2511	
SEQ2501	ACTTAT-AGTTCTGAGACTAATCAAACAACTCATCAAAATTACTATAATAGTAGCACTC
SEQ2502	ACTTAT-AGTTCTGAGACTAATCAAACAACTCATCAAAGTTACTATAATAGTAGCACTC
SEQ2503	ACTTAT-AGTTCTGAGACTAATCAAACAACTCATCAAAATTACTATAATAGTAGCACTC
SEQ2504	ACTTAT-AGTTCTGAGACTAATCAAACAACTCATCAAAATTACTATAATAGTAGCACTC
SEQ2505	ACTTAT-AGTTCTGAGACTAATCAAACAACTCATCAAAATTACTATAATAGTAGCACTC
SEQ2506	ACTTATTAGTTCTGAGACTAATCAAACAACTCATCAAAATTACTATAATAGTAGCACTC
SEQ2507	
SEQ2508	
SEQ2509	
SEQ2509 SEQ2510	ACTTAT-AGTTCTGAGACTAATCAAACAACTCATCAAAATTACTATAATAGTAGCACTC
SEQ2511	

## Table 25: Comparative Sequences relating t SAG0368 (protein of unknown functi n)

SEQ2501	TGCTAGTAACTATAGCAGTAACACTAACACAGGTCAGGCTGATTCAAGTGGAAGTGTCA
SEQ2502	TGCTAATAACTATAGCAGTAACACTAACACAGGTCAGGCTGATTCAAGTGGAAGTGTCA
SEQ2503	TGCTAGTAACTATAGCAGTAACACTAACACAGGTCAGGCTGATTCAAGTGGAAGTGTCA
SEQ2504	TGCTAGTAACTATAGCAGTAACACTAACACGGTCAGGCTGATTCAAGTGGAAGTGTCA
SEQ2505	TGCTAGTAACTATAGCAGTAACACTAACACAGGTCAGGCTGATTCAAGTGGAAGTGTCA
SEQ2506	TGCTAGTAACTATAGCAGTAACACTAACACAGGTCAGGCTGATTCAAGTGGAAGTGTCA
SEQ2507	
SEQ2508	
SEQ2509	
SEQ2510	TGCTAGTAACTATAGCAGTAACACTAACACAGGTCAGGCTGATTCAAGTGGAAGTGTCA
SEQ2511	
SEQ2501	TAATCATAACGGGGCTGCAACGCCTAATCCA
SEQ2502	TAATCATAATGGGGCTGCAACGCCTAATCCA
SEQ2503	TAATCATAACGGGGCTGCAACGCCTAATCCA
SEQ2504	TAATCATAACGGGGCTGCAACGCCTAATCCA
SEQ2505	TAATCATAACGGGGCTGCAACGCCTAATCCA
SEQ2506	TAATCATAACGGGGCTGCAACGCCTAATCCA
SEQ2507	00440
SEQ2508	
SEQ2509	
· SEQ2510	TAATCATAACGGGGCTGCAACGCCTAATCCA
SEQ2511	***************************

#### >SEQ ID NO 2550: 54 090 frame: 1

YNFSTNELSKTFKDFKLAKSKSHAIEETKPFSILLMGVDTGSEHRKSKWSGNSDSMILVT INPKTNKTTMTSLERDVLIKLSGPKNNGQTGVEAKLNAAYASGGAEMALMTVQDLLDINV DYFMQINMQGLVDLVNAVGGITVTNKFDFPISIAANEPEYKAVVEPGTHKINGEQALVYS RMRYDDPEGDYGRQKRQREVIQKVLKKILALNSISSYKKILSAVSNNMQTNIEISSKTIP NLLAYKDSLEHIKSYQLKGEDATLSDGGSYQILTKKHLLAVQNRIKKELDKKRSKTLKTS AILYEDYYGTTASNDSSTYSSTQENNYNTTPYSEAPPSYSGNTTYSSETNQTTHQNYYNS STPASNYSSNTNTGOADSSGSVNNHNGAATPNP

#### >SEQ ID NO 2551:54 1169NT frame: 1

YNFSTNELSKTFKDFKLAKSKSHAIEETKPFSILLMGVDTGSEHRKSKLVRK.RFYDLSH
YKS.N..NNDDKLRT.RID.IEWSQK.WTDWRRSKAKCSLCFWWCGNGIDDCSRLIRY.C
.LLYAN.YARIS.FSQCCWWYNSN..I.LSNINCCQ.TRVQGCC.TRDT.NKWRTSTCLF
SYAL..SRGRLWASKKTT.SNSKSP.KNIGVK.Y.FIQKNSFRSK..HAN.Y.DIIKNDS
.FVSL.RFIGTY.ILSVER.RRYFIRWWLLSNFN.ETSTCSSK.N.ERTR.KA..NSEDK
RDSI.RLLWYYC...FFYLFINTRE.L.YNTLFRSTTKLQW.YYL.F.D.SNNSSKLL..
.HSC..L.Q.H.HRSG.FKWKCQ.S.WGCNA.S

#### >SEQ ID NO 2552:54\_18RS21 frame: 1

YNFSTNELSKTFKDFKLAKSKSHAIEETKPFSILLMGVDTGSEHRKSKWSGNSDSMILVT INPKTNKTTMTSLERDVLIKLSGPKNNGQTGVEAKLNAAYASGGAEMALMTVQDLLDINV DYFMQINMQGLVDLVNAVGGITVTNKFDFPISIAANEPEYKAVVEPGTHKINGEQALVYS RMRYDDPEGDYGRQKRQREVIQKVLKKILALNSISSYKKILSAVSNNMQTNIEISSKTIP NLLAYKDSLEHIKSYQLKGEDATLSDGGSYQILTKKHLLAVQNRIKKELDKKRSKTLKTS AILYEDYYGTTASNDSSTYSSTQENNYNTTPYSEAPPSYSGNTTYSSETNQTTHQNYYNS STPASNYSSNTNTGQADSSGSVNNHNGAATPNP

#### >SEQ ID NO 2553:54 2603 frame: 1

YNFSTNELSKTFKDFKLAKSKSHAIEETKPFSILLMGVDTGSEHRKSKWSGNSDSMILVT
INPKTNKTTMTSLERDVLIKLSGPKNNGQTGVEAKLNAAYASGGAEMALMTVQDLLDINV
DYFMQINMQGLVDLVNAVGGITVTNKFDFPISIAANEPEYKAVVEPGTHKINGEQALVYS
RMRYDDPEGDYGRQKRQREVIQKVLKKILALNSISSYKKILSAVSNNMQTNIEISSKTIP
NLLAYKDSLEHIKSYQLKGEDATLSDGGSYQILTKKHLLAVQNRIKKELDKKRSKTLKTS
AILYEDYYGTTASNDSSTYSSTQENNYNTTPYSEAPPSYSGNTTYSSETNQTTHQNYYNS
STPASNYSSNTNTGQADSSGSVNNHNGAATPNP

#### >SEQ ID NO 2554: 54\_A909 frame: 1

YNFSTNELSKTFKDFKLAKSKSHAIEETKPFSILLMGVDTGSEHRKSKWSGNSDSMILVT INPKTNKTTMTSLERDVLIKLSGPKNNGQTGVEAKLNAAYASGGAEMALMTVQDLLDINV DYFMQINMQGLVDLVNAVGGITVTNKFDFPISIAANEPEYKAVVEPGTHKINGEQALVYS RMRYDDPEGDYGRQKRQREVIQKVLKKILALNSISSYKKILSAVSNNMQTNIEISSKTIP NLLAYKDSLEHIKSYQLKGEDATLSDGGSYQILTKKHLLAVQNRIKKELDKKRSKTLKTS



AILYEDYYGTTASNDSSTYSSTQENNYNTTPYSEAPPSYSGNTTYSSETNQTTHQNYYNS STPASNYSSNTNTGOADSSGSVNNHNGAATPNP

#### >SEQ ID NO 2555:54 CJB110 frame: 1

YNFSTNELSKTFKDFKLAKSKSHAIEETKPFSILLMGVDTGSEHRKSKWSGNSDSMILVT INPKTNKTTMTSLERDVLIKLSGPKNNGQTGVEAKLNAAYASGGAEMALMTVQDLLDINV DYFMQINMQGLVDLVNAVGGITVTNKFDFPISIAANEPEYKAVVEPGTHKINGEQALVYS RMRYDDPEGDYGRQKRQREVIQKVLKKILALNSISSYKKILSAVSNNMQTNIEISSKTIP NLLAYKDSLEHIKSYQLKGEDATLSDGGSYQILTKKHLLAVQNRIKKELDKKRSKTLKTS AILYEDYYGTTASNDSSTYSSTQENNYNTTPYSEAPPSYSGNTTY.F.D.SNNSSKLL..

#### >SEQ ID NO 2556:54\_COH1 frame: 1

DFKLDKSKSHAIEETKPFSILLMGVDTGSEHRKSKWSGNSDSMILVTINPKTNKTTMTSL ERDVLIKLSGPKNNGQTGVEAKLNAAYASGGAEMALMTVQDLLDINVDYFMQINMQGLVD LVNAVGGITVTNKFDFPISIAANEPEYKAVVEPGTHKINGEQALVYSRMRYDDPEGDYGR QKRQREVIQKVLKKILALNSISSYKKILSAVSNNMQTNIEISSKTIPNLLAYKDSLEHIK SYQLKGEDATLSDGGSYQILTKKHLLAVQNRIKKELDKKRSKTLKTSAILYEDYYGTTAS NDSSTYSSTQENYYYTTPLFRSTTKLQW.YYL.F.D.SNNSSKLL...HSC..L.Q.H.H RSG.FKWKC..L.RGCNA.SKHRNATSTRSN.S

#### >SEQ ID NO 2557:54 H36B frame: 1

YNFSTNELSKTFKDFKLAKSKSHAIEETKPFSILLMGVDTGSEHRKSKWSGNSDSMILVT INPKTNKTTMTSLERDVLIKLSGPKNNGQTGVEAKLNAAYASGGAEMALMTVQDLLDINV DYFMQINMQGLVDLVNAVGGITVTNKFDFPISIAANEPEYKAVVEPGTHKINGEQALVYS RMRYDDPEGDYGRQKRQREVIQKVLKKILALNSISSYKKILSAVSNNMQTNIEISSKTIP NLLAYKDSLEHIKSYQLKGEDATLSDGGSYQILTKKHLLAVQNRIKKELDKKRSKTLKTS AILYEDYYGTTASNDSSTYSSTQENNYNTTPYSEAPPSYSGNTTYSSETNQTTHQNYYNS STPASNYSSNTNTGQADSSGSVNNHNGAATPNP

#### >SEQ ID NO 2558:54 JM9130013 frame: 1

YNFSTNELSKTFKDFKLAKSKSHAIEETKPFSILLMGVDTGSEHRKSKWSGNSDSMILVT INPKTNKTTMTSLERDVLIKLSGPKNNGQTGVEAKLNAAYASGGAEMALMTVQDLLDINV DYFMQINMQGLVDLVNAVGGITVTNKFDFPISIAANEPEYKAVVEPGTHKINGEQALVYS RMRYDDPEGDYGRQKRQREVIQKVLKKILALNSISSYKKILSAVSNNMQTNIEISSKTIP NLLAYKDSLEHIKSYQLKGEDATLSDGGSYQILTKKHLLAVQNRIKKELDKKRSKTLKTS AILYEDYYGTTASNDSSTYSSTQENNYNTTPYSEAPPSYSGNTTYSSETNQTTHQNYYNS STPASNYSSNTNTGQADSSGSVNNHNGAATPNP

#### >SEQ ID NO 2559:54 M781 frame: 2

SILLMGVDTGSEHRKSKWSGNSDSMILVTINPKTNKTTMTSLERDVLIKLSGPKNNGQTG
VEAKLNAAYASGAEMALMTVQDLLDINVDYFMQINMQGLVDLVNAVGGITVTNKFDFPI
SIAANEPEYKAVVEPGTHKINGEQALVYSRMRYDDPEGDYGRQKRQREVIQKVLKKILAL
NSISSYKKILSAVSNNMQTNIEISSKTIPNLLAYKDSLEHIKSYQLKGEDATLSDGGSYQ
ILTKKHLLAVQNRIKKELDKKRSKTLKTSAILYEDYYGTTASNDSSTYSSTQENNYNTTP
YSEAPPSYSGNTTYSSETNQTTHQSYYNSSTPASNYSSNTNTGQADSSGSVNNYNGAATP
NPNTGTQPVPGQTNP

SEQ2550	NFSTNELSKTFKDFKLAKSKSHAIEETKPFSILLMGVDTGSEHRKSKWSGNSDSMILVT
SEQ2551	NFSTNELSKTFKDFKLAKSKSHAIEETKPFSILLMGVDTGSEHRKSKLVRKRFYDLSHY
SEQ2552	NESTNELSKTFKDFKLAKSKSHAIEETKPFSILLMGVDTGSEHRKSKWSGNSDSMILVT
SEQ2553	NFSTNELSKTFKDFKLAKSKSHAIEETKPFSILLMGVDTGSEHRKSKWSGNSDSMILVT
SEQ2554	NFSTNELSKTFKDFKLAKSKSHAIEETKPFSILLMGVDTGSEHRKSKWSGNSDSMILVT
SEQ2555	NFSTNELSKTFKDFKLAKSKSHAIEETKPFSILLMGVDTGSEHRKSKWSGNSDSMILVT
SEQ2556	DFKLDKSKSHAIEETKPFSILLMGVDTGSEHRKSKWSGNSDSMILVT
SEQ2557	NFSTNELSKTFKDFKLAKSKSHAIEETKPFSILLMGVDTGSEHRKSKWSGNSDSMILVT
SEQ2558	NFSTNELSKTFKDFKLAKSKSHAIEETKPFSILLMGVDTGSEHRKSKWSGNSDSMILVT
SEQ2559	SILLMGVDTGSEHRKSKWSGNSDSMILVT
SEQ2550	NPKTNKTTMTSLERDVLIKLSGPKNNGOTGVEAKLNAAYASGGAEMALMTVQDLLDINV
SEQ2551	SNNNDDKLRTRIDIEWSOKWTDWRRSKAKCSLCFWWCGNGIDDCSRLIRYCLLY
SEQ2552	NPKTNKTTMTSLERDVLIKLSGPKNNGOTGVEAKLNAAYASGGAEMALMTVQDLLDINV
SEQ2553	NPKTNKTTMTSLERDVLIKLSGPKNNGOTGVEAKLNAAYASGGAEMALMTVQDLLDINV
SEQ2554	NPKTNKTTMTSLERDVLIKLSGPKNNGOTGVEAKLNAAYASGGAEMALMTVQDLLDINV
SEQ2555	NPKTNKTTMTSLERDVLIKLSGPKNNGOTGVEAKLNAAYASGGAEMALMTVQDLLDINV
SEQ2556	NPKTNKTTMTSLERDVLIKLSGPKNNGOTGVEAKLNAAYASGGAEMALMTVQDLLDINV
SEQ2557 ·	NPKTNKTTMTSLERDVLIKLSGPKNNGQTGVEAKLNAAYASGGAEMALMTVQDLLDINV



SEQ2558	npktnkttmtslerdvliklsgpknngQtgveaklnaayasggaemalmtvQdlldinv
SEQ2559	NPKTNKTTMTSLERDVLIKLSGPKNNGQTGVEAKLNAAYASGGAEMALMTVQDLLDINV
	<del>-</del>
SEQ2550	YFMQINMQGLVDLVNAVGGITVTNKFDFPISIAANEPEYKAVVEPGTHKINGEQALVYS
SEQ2551	NYARISFSQCCWWYNSNILSNINCCQTRVQGCCTRDTNKWRTSTCLFSY
SEQ2552	YFMQINMQGLVDLVNAVGGITVTNKFDFPISIAANEPEYKAVVEPGTHKINGEQALVYS
SEQ2553	YFMQINMQGLVDLVNAVGGITVTNKFDFPISIAANEPEYKAVVEPGTHKINGEQALVYS
SEQ2554	YFMQINMQGLVDLVNAVGGITVTNKFDFPISIAANEPEYKAVVEPGTHKINGEOALVYS
SEQ2555	YFMQINMQGLVDLVNAVGGITVTNKFDFPISIAANEPEYKAVVEPGTHKINGEQALVYS
SEQ2556	YFMQINMQGLVDLVNAVGGITVTNKFDFPISIAANEPEYKAVVEPGTHKINGEQALVIS
SEQ2557	YFMQINMQGLVDLVNAVGGITVTNKFDFPISIAANEPEYKAVVEPGTHKINGEQALVYS
SEQ2558	TENGINAGGIVDLVNAVGGIIVINAFDFFISAANEPEIRAVVEPGTHKINGEQALVYS
_	YFMQINMQGLVDLVNAVGGITVTNKFDFPISIAANEPEYKAVVEPGTHKINGEQALVYS
SEQ2559	YFMQINMQGLVDLVNAVGGITVTNKFDFPISIAANEPEYKAVVEPGTHKINGEQALVYS
SEQ2550	MRYDDPEGDYGRQKRQREVIQKVLKKILALNSISSYKKILSAVSNNMQTNIEISSKTIP
SEQ2551	LSRGRLWASKKTTSNSKSPKNIGVKYFIQKNSFRSKHANYDIIKNDSFVSLRFIGTYI-
SEQ2552	MRYDDPEGDYGRQKRQREVIQKVLKKILALNSISSYKKILSAVSNNMQTNIEISSKTIP
SEQ2553	MRYDDPEGDYGRQKRQREVIQKVLKKILALNSISSYKKILSAVSNNMQTNIEISSKTIP
SEQ2554	MRYDDPEGDYGRQKRQREVIQKVLKKILALNSISSYKKILSAVSNNMQTNIEISSKTIP
SEQ2555	MRYDDPEGDYGRQKRQREVIQKVLKKILALNSISSYKKILSAVSNNMQTNIEISSKTIP
SEQ2556	MRYDDPEGDYGRQKRQREVIQKVLKKILALNSISSYKKILSAVSNNMQTNIEISSKTIP
SEQ2557	MRYDDPEGDYGRQKRQREVIQKVLKKILALNSISSYKKILSAVSNNMQTNIEISSKTIP
SEQ2558	MRYDDPEGDYGRQKRQREVIQKVLKKILALNSISSYKKILSAVSNNMOTNIEISSKTIP
SEQ2559	MRYDDPEGDYGRQKRQREVIQKVLKKILALNSISSYKKILSAVSNNMQTNIEISSKTIP
_	
SEQ2550	LLAYKDSLEHIKSYQLKGEDATLSDGGSYQILTKKHLLAVQNRIKKELDKKRSKTLKTS
SEQ2551	L-SVERRRYFIRWWLLSNFNETSTCSSKNERTRKANSEDKRDSIRLLWYYCFFYLFINT
SE02552	LLAYKDSLEHIKSYQLKGEDATLSDGGSYQILTKKHLLAVONRIKKELDKKRSKTLKTS
SEQ2553	LLAYKDSLEHIKSYQLKGEDATLSDGGSYQILTKKHLLAVQNRIKKELDKKRSKTLKTS
SEQ2554	LLAYKDSLEHIKSYQLKGEDATLSDGGSYQILTKKHLLAVONRIKKELDKKRSKTLKTS
SEQ2555	LLAYKDSLEHIKSYQLKGEDATLSDGGSYQILTKKHLLAVQNRIKKELDKKRSKTLKTS
SEQ2556	LLAYKOSLEHIKSYQLKGEDATLSDGGSYQILTKKHLLAVQNRIKKELDKKRSKTLKTS
SEQ2557	LLAYKDSLEHIKSYQLKGEDATLSDGGSYQILTKKHLLAVQNRIKKELDKKRSKTLKTS
SEQ2558	LLAYKDSLEHIKSYQLKGEDATLSDGGSYQILTKKHLLAVQNRIKKELDKKRSKTLKTS
SEQ2559	
5602333	LLAYKDSLEHIKSYQLKGEDATLSDGGSYQILTKKHLLAVQNRIKKELDKKRSKTLKTS
SEQ2550	TY VID VICENTA OVER COMPANIES OF THE PROPERTY
SEQ2551	ILYEDYYGTTASNDSSTYSSTQENNYNTTPYSEAPPSYSGNTTYSSETNQTTHQNYYNS
	ELYTKLQWYYLFDSNNSSKLLHSCLQH
SEQ2552	ILYEDYYGTTASNDSSTYSSTQENNYNTTPYSEAPPSYSGNTTYSSETNQTTHQNYYNS
SEQ2553	ILYEDYYGTTASNDSSTYSSTQENNYNTTPYSEAPPSYSGNTTYSSETNQTTHQNYYNS
SEQ2554	ILYEDYYGTTASNDSSTYSSTQENNYNTTPYSEAPPSYSGNTTYSSETNQTTHQNYYNS
SEQ2555	ILYEDYYGTTASNDSSTYSSTQENNYNTTPYSEAPPSYSGNTTYFDSNNSSKLL
SEQ2556	ILYEDYYGTTASNDSSTYSSTQENYYYTTPLFRSTTKLQWYYLFDSNNSSKLLHSCLQH
SEQ2557	ILYEDYYGTTASNDSSTYSSTQENNYNTTPYSEAPPSYSGNTTYSSETNQTTHQNYYNS
SEQ2558	ILYEDYYGTTASNDSSTYSSTQENNYNTTPYSEAPPSYSGNTTYSSETNQTTHQNYYNS
SEQ2559	ILYEDYYGTTASNDSSTYSSTQENNYNTTPYSEAPPSYSGNTTYSSETNQTTHQSYYNS
SEQ2550	TPASNYSSNTNTGQADSSGSVNNHNGAATPNP
SEQ2551	RSGFKWKCQSWGCNAS
SEQ2552	TPASNYSSNTNTGQADSSGSVNNHNGAATPNP
SEQ2553	TPASNYSSNTNTGQADSSGSVNNHNGAATPNP
SEQ2554	TPASNYSSNTNTGQADSSGSVNNHNGAATPNP
SEQ2555	***************************************
SEQ2556	RSGFKWKCLRGCNASKHRNATSTRSNS
SEQ2557	TPASNYSSNTNTGQADSSGSVNNHNGAATPNP
SEQ2558	TPASNYSSNTNTGQADSSGSVNNHNGAATPNP
SEQ2559	TPASNYSSNTNTGQADSSGSVNNYNGAATPNPNTGTQPVPGQTNP
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#### Table 26: C mparative Sequences relating to SAG0503 (lipase/acylhydolase)

## SECTIO NO. 2601: SAG0503 FROM THE 090 GBS TYPE IA STRAIN (REVERSE COMPLEMENT)

## SEQ ID NO. 2602: SAG0503 FROM THE H36b GBS TYPE Ib STRAIN (REVERSE COMPLEMENT)

### SEQ ID NO. 2603: SAG0503 FROM THE 18RS21 GBS TYPE II STRAIN (REVERSE COMPLEMENT)

## SEQ ID NO. 2604: SAG0503 FROM THE COH1 GBS TYPE III STRAIN (REVERSE COMPLEMENT)

#### Table 26: Comparative Sequences relating t SAG0503 (lipase/acylhyd lase)

## SEC\_ID NO. 2605: SAG0503 FROM THE CJB110 GBS NONTYPEABLE STRAIN (REVERSE COMPLEMENT)

### SEQ ID NO. 2606: SAG0503 FROM THE 1169NT1 GBS TYPE V STRAIN (REVERSE COMPLEMENT)

## SEQ ID NO. 2607: SAG0503 FROM THE JM9130013 GBS TYPE VIII STRAIN (REVERSE COMPLEMENT)

## SEQ ID NO. 2608: SAG0503 FROM THE 2603 V/R GBS TYPE V STRAIN (REVERSE COMPLEMENT)

### Table 26: C mparative Sequences relating t SAG0503 (lipase/acylhydolase)

SEC\_ID NO. 2609: SAG0503 FROM THE M781 CBS TYPE III STRAIN (REVERSE COMPLEMENT)

GGACAAGTTTGTACAAAAAAGCAGGCTCTATTTTTTCCTTGATCATTCCAAAATCAAATCCTAAATTAACAAAAAAAG ACTTCCTAACAAAGAAAGTTATCCCACTTAACTATGTTGCTCTTGGAGATTCTCTGACCGAAGGTGTGGGGGATACAA CCTCTCAAGGTGGTTTTGTCCCACTGCTATCAGAATCACTCCATAATCGATACTCTTACCAAGTGACTTCTGTTAATT ATGGTGTGTCTGGGAATACTAGTCAACAATTTTAAAACGTATGACGACAGATCCTCAAATCGAAAAAGATTTAGAGA  ${\tt AAGCTGATTTATTGACGCTAACTGTTGGTGGTAATGATGTCTTGGCTGTTATTCGTAAAGAGCTCAGTCATTTATCAC}$ CTAAATTGCCTATTTATGTTTTAGGCATTTATAATCCTTTTTACCTAAACTTTCCACAATTAACTAAAATGCAAACCG TTATTGATAATTGGAATAAAGCTACAAAAGAAGTAGTTGATGCTTCAGAAAATGTTTATTTTGTCCCAATTAATGACC GCCTTTATAAGGGAATAAATGGTAAAGAGGGTATTACAGAGTCATCAAATAGTCAGGCAAGTATCACTAATGATGCTC AAACAAGAAAAACTGGCCGAACCCAGCTTTCTTGTACAAA

SEQ2601	GGCACAAGTTTGTACAAAAAAGCAGGCTCTATTTTTTCCTTGATCATTCCAAAATCAAA
SEQ2602	TTTGTACAAAAAGCAGGCTCTATTTTTTCCTTGATCATTCCAAAATCAAA
SEQ2603	GTTTGTACAAAAAGCAGGCTCTATTTTTCCTTGATCATTCCAAAATCAAA
SEQ2604	GGACAAGTTTGTACAAAAAAGCAGGCTCTATTTTTTCCTTGATCATTCCAAAATCAAA
SEQ2605	GTTTGTACAAAAAGCAGGCTCTATTTTTCCTTGATCATTCCAAAATCAAA
SEQ2606	GTTTGTACAAAAAGCAGGCTCTATTTTTCCTTGATCATTCCAAAATCAAA
SEQ2607	GTTTGTACAAAAAGCAGGCTCTATTTTTTCCTTGATCATTCCAAAATCAAA
SEQ2608	AGTTTGTACAAAAAGCAGGCTCTATTTTTTCCTTGATCATTCCAAAATCAAA
SEQ2609	GGACAAGTTTGTACAAAAAAGCAGGCTCTATTTTTTCCTTGATCATTCCAAAATCAAA
SEQ2601	TCCTAAATTAACAAAAAAAGACTTCCTAACAAAGAAAGTTATCCCACTTAACTATGTTGC
SEQ2602	TCCTAAATTAACAAAAAAAGACTTCCTAACAAAGAAAGTTATCCCACTTAACTATGTTGC
SEQ2603	TCCTAAATTAACAAAAAAAGACTTCCTAACAAAGAAAGTTATCCCACTTAACTATGTTGC
SEQ2604	TCCTAAATTAACAAAAAAAGACTTCCTAACAAAGAAAGTTATCCCACTTAACTATGTTGC
SEQ2605	TCCTAAATTAACAAAAAAAGACTTCCTAACAAAGAAAGTTATCCCACTTAACTATGTTGC
SEQ2606	TCCTAAATTAACAAAAAAAGACTTCCTAACAAAGAAAGTTATCCCACTTAACTATGTTGC
SEQ2607	TCCTAAATTAACAAAAAAGCTTCCTAACAAGAAAGTTATCCCACTTAACTATGTTGC
SEQ2608	TCCTAAATTAACAAAAAAAGCTTCCTAACAAAGAAAGTTATCCCACTTAACTATGTTGC
SE02609	TCCTAAATTAACAAAAAAAGCTTCCTAACAAAGAAAGTTATCCCACTTAACTATGTTGC
•	, , , , , , , , , , , , , , , , , , ,
SEQ2601	TCTTGGAGATTCTCTGACCGAAGGTGTGGGCGATACAACCTCTCAAGGTGGTTTTGTCCC
SEQ2602	TCTTGGAGATTCTCTGACCGAAGGTGTGGGCGATACAACCTCTCAAGGTGGTTTTGTTCC
SEQ2603	TCTTGGAGATTCTCTGACCGAAGGTGTGGGCGATACAACCTCTCAAGGTGGTTTTGTTCC
SEQ2604	TCTTGGAGATTCTCTGACCGAAGGTGTGGGGGGATACAACCTCTCAAGGTGGTTTTGTCCC
SEQ2605	TCTTGGAGATCTCTGACCGAAGGTGTGGGCGATACAACCTCTCAAGGTGGTTTTGTCCC
SEQ2606	TCTTGGAGATTCTCTGACCGAAGGTGTGGGGGGATACAACCTCTCAAGGTGGTTTTGTCCC
SEQ2607	TCTTGGAGATCTCTGACCGAAGGTGTGGGCGATACAACCTCTCAAGGTGGTTTTGTCCC
SEQ2608	TCTTGGAGATTCTCTGACCGAAGGTGTGGGCGATACAACCTCTCAAGGTGGTTTTGTTCC
SEQ2609	TCTTGGAGATTCTCTGACCGAAGGTGTGGGGGGATACAACCTCTCAAGGTGGTTTTGTCCC
	TOTTO CONTROL OF THE PROPERTY
SEQ2601	ACTGCTATCAGAATCACTCCATAATCGATACTCTTACCAAGTGACTTCTGTTAATTATGG
SEQ2602	ACTGCTATCAGAATCACTCCATAATCGATACTCTTACCAAGTGACTTCTGTTAATTATGG
SEQ2603	ACTGCTATCAGAATCACTCCATAATCGATACTCTTACCAAGTGACTTCTGTTAATTATGG
SEQ2604	ACTGCTATCAGAATCACTCCATAATCGATACTCTTACCAAGTGACTTCTGTTAATTATGG
SEQ2605	ACTGCTATCAGAATCACTCCATAATCGATACTCTTACCAAGTGACTTCTGTTAATTATGG
SEQ2606	ACTGCTATCAGAATCACTCCATAATCGATACTCTTACCAAGTGACTTCTGTTAATTATGG
SEQ2607	ACTGCTATCAGAATCACTCCATAATCGATACTCTTACCAAGTGACTTCTGTTAATTATGG
SEQ2608	ACTGCTATCAGAATCACTCCATAATCGATACTCTTACCAAGTGACTTCTGTTAATTATGG
SEQ2609	ACTGCTATCAGAATCACTCCATAATCGATACTCTTACCAAGTGACTTCTGTTAATTATGG
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SEQ2601	TGTGTCTGGGAATACTAGTCAACAAATTTTAAAACGTATGACGACAGATCCTCAAATCGA
SEQ2602	TGTGTCTGGGAATACTAGTCAACAAATTTTAAAACGTATGACGACAGATCCTCAAATCGA
SEQ2603	TGTGTCTGGGAATACTAGTCAACAAATTTTAAAACGTATGACGACAGATCCTCAAATCGA
SEQ2604	TGTGTCTGGGAATACTAGTCAACAAATTTTAAAACGTATGACGACAGATCCTCAAATCGA
SEQ2605	TGTGTCTGGGAATACTAGTCAACAAATTTTAAAACGTATGACGACAGATCCTCAAATCGA
SEQ2606	TGTGTCTGGGAATACTAGTCAACAAATTTTAAAACGTATGACGACAGATCCTCAAATCGA
SEQ2607	TGTGTCTGGGAATACTAGTCAACAAATTTTAAAACGTATGACGACAGATCCTCAAATCGA
SEQ2608	TGTGTCTGGGAATACTAGTCAACAAATTTTAAAACGTATGACGACAGATCCTCAAATCGA
SEQ2609	TGTGTCTGGGAATACTAGTCAACAAATTTTAAAACGTATGACGACAGATCCTCAAATCGA TGTGTCTGGGAATACTAGTCAACAAATTTTAAAACGTATGACGACAGATCCTCAAATCGA
SEQ2601	AAAAGATTTAGAGAAAGCTGATTTATTGACGCTAACTGTTGGTGGTAATGATGTCTTGGC
SEQ2602	AAAAGATTTAGAGAAAGCTGATTTATTGACGCTAACTGTTGGTGGTAATGATGTCTTTGGC AAAAGATTTAGAGAAAGCTGATTTATTGACGCTAACTGTTGGTGGTAATGATGTCTTTGGC
SEQ2603	AAAAGATTTAGAGAAAGCTGATTTATTGACGCTAACTGTTGGTGGTAATGATGTCTTGGC AAAAGATTTAGAGAAAGCTGATTTATTGACGCTAACTGTTGGTGGTAATGATGTCTTGGC
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SEC. 304	AAAAGATTTAGAGAAAGCTGATTTATTGACGCTAACTGTTGGTGGTAATGATGTCTTGGC
SEQ2605	AAAAGATTTAGAGAAAGCTGATTTATTGACGCTAACTGTTGGTGGTAATGATGTCTTGGC
SEQ2606	AAAAGATTTAGAGAAAAGCTGATTTATTGACGCTAACTGTTGGTGGTAATGATGTCTTGGC
SEQ2607 SEQ2608	AAAAGATTTAGAGAAAGCTGATTTATTGACGCTAACTGTTGGTGGTAATGATGTCTTGGC
SEQ2609	AAAAGATTTAGAGAAAGCTGATTTATTGACGCTAACTGTTGGTGGTAATGATGTCTTGGC
250203	AAAAGATTTAGAGAAAGCTGATTTATTGACGCTAACTGTTGGTGGTAATGATGTCTTGGC
SEQ2601	TGTTATTCGTAAAGAGCTCAGTCATTTATCACTAAATTCCTTTGAGAAACCAGCAGAAGC
SEQ2602	TGTTATTCGTAAAGAGCTCAGTCATTTATCACTAAATTCCTTTGAGAAACCAGCAGAAGC
SEQ2603	TGTTATTCGTAAAGAGCTCAGTCATTTATCACTAAATTCCTTTGAGAAACCAGCAGAAGC
SEQ2604	TGTTATTCGTAAAGAGCTCAGTCATTTATCACTAAATTCCTTTGAGAAACCAGCAGAAGC
SEQ2605	TGTTATTCGTAAAGAGCTCAGTCATTTATCACTAAATTCCTTTGAGAAACCAGCAGAAGC
SEQ2606	TGTTATTCGTAAAGAGCTCAGTCATTTATCACTAAATTCCTTTGAGAAACCAGCAGAAGC
SEQ2607	TGTTATTCGTAAAGAGCTCAGTCATTTATCACTAAATTCCTTTGAGAAACCAGCAGAAGC
SEQ2608	TGTTATTCGTAAAGAGCTCAGTCATTTATCACTAAATTCCTTTGAGAAACCAGCAGAAGC
SEQ2609	TGTTATTCGTAAAGAGCTCAGTCATTTATCACTAAATTCCTTTGAGAAACCAGCAGAAGC
GT-00-C01	101011 CO11 COUNTY 101 101 101 101 101 101 101 101 101 10
SEQ2601	ATATAAGGAACGTTTGAAAGAAATACTTGCAAAAGCAAGACAAGATAATCCTAAATTGCC
SEQ2602	ATATAAGGAACGTTTGAAAGAAATCCTTGCAAAAGCAAGACAAGATAATCCCTAAATTGCC
SEQ2603 SEQ2604	ATATAAGGAACGTTTGAAAGAAATCCTTGCAAAAGCAAGACAAGATAATCCTAAATTGCC
SEQ2604 SEQ2605	ATATAAGGAACGTTTGAAAGAAATTCTTGCAAAAGCAAGACAAGATAATCCTAAATTGCC
SEQ2606	ATATAAGGAACGTTTGAAAGAAATACTTGCAAAAGCAAGACAAGATAATCCTAAATTGCC ATATAAGGAACGTTTGAAAGAAATTCTTGCAAAAGCAAGACAAGATAATCCTAAATTGCC
SEQ2607	ATATAAGGAACGTTTGAAAGAAATCCTTGCAAAAGCAAGACAAGATAATCCTAAATTGCC ATATAAGGAACGTTTGAAAGAAATCCTTGCAAAAGCAAGACAAGATAATCCTAAATTGCC
SEQ2608	ATATAAGGAACGTTTGAAAGAAATCCTTGCAAAAGCAAGACAAGATAATCCTAAATTGCC ATATAAGGAACGTTTGAAAGAAATCCTTGCAAAAGCAAGACAAGATAATCCTAAATTGCC
SEQ2609	ATATAAGGAACGTTTGAAAGAAATTCTTGCAAAAGCAAGACAAGATAATCCTAAATTGCC
0282003	ATATAMAGAACGTTTGAAAAAATTGCC
SEQ2601	TATTTATGTTTTAGGCATTTATAATCCTTTTTACCTAAACTTTCCACAATTAACTAAAAT
SEQ2602	TATTTATGTTTTAGGCATTTATAATCCTTTTTACCTAAACTTTCCACAATTAACTAAAAT
SEQ2603	TATTTATGTTTTAGGCATTTATAATCCTTTTTACCTAAACTTTCCACAATTAACTAAAAT
SEQ2604	TATTTATGTTTTAGGCATTTATAATCCTTTTTACCTAAACTTTCCACAATTAACTAAAAT
SEQ2605	TATTTATGTTTTAGGCATTTATAATCCTTTTTACCTAAACTTTCCACAATTAACTAAAAT
SEQ2606	TATTTATGTTTTAGGCATTTATAATCCTTTTTACCTAAACTTTCCACAATTAACTAAAAT
SEQ2607	TATTTATGTTTTAGGCATTTATAATCCTTTTTACCTAAACTTTCCACAATTAACTAAAAT
SEQ2608	TATTTATGTTTTAGGCATTTATAATCCTTTTTACCTAAACTTTCCACAATTAACTAAAAT
SEQ2609	TATTTATGTTTTAGGCATTTATAATCCTTTTTACCTAAACTTTCCACAATTAACTAAAAT
SEQ2601	GCAAACCGTTATTGATAATTGGAATAAAGCTACAAAAGAAGTAGTTGATGCTTCAGAAAA
SEQ2602	GCAAACCGTTATTGATAATTGGAATAAAGCTACAAAAGAAGTAGTTGATGCTTCAGAAAA
SEQ2603	GCAAACCGTTATTGATAATTGGAATAAAGCTACAAAAGAAGTAGTTGATGCTTCAGAAAA
SEQ2604	GCAAACCGTTATTGATAATTGGAATAAAGCTACAAAAGAAGTAGTTGATGCTTCAGAAAA
SEQ2605	GCAAACCGTTATTGATAATTGGAATAAAGCTACAAAAGAAGTAGTTGATGCTTCAGAAAA
SEQ2606	GCAAACCGTTATTGATAATTGGAATAAAGCTACAAAAGAAGTAGTTGATGCTTCAGAAAA
SEQ2607	GCAAACCGTTATTGATAATTGGAATAAAGCTACAAAAGAAGTAGTTGATGCTTCAGAAAA
SEQ2608	GCAAACCGTTATTGATAATTGGAATAAAGCTACAAAAGAAGTAGTTGATGCTTCAGAAAA
SEQ2609	GCAAACCGTTATTGATAATTGGAATAAAGCTACAAAAGAAGTAGTTGATGCTTCAGAAAA
CP02601	
SEQ2601	TGTTTATTTTGTCCCAATTAATGACCGCCTTTATAAGGGAATAAATGGTAAAGAGGGTAT
SEQ2602 SEQ2603	TGTTTATTTTGTCCCAATTAATGACCGCCTTTATAAGGGAATAAATGGTAAAGAGGGTAT
SEQ2604	TGTTTATTTTGTCCCAATTAATGACCGCCTTTATAAGGGAATAAATGGTAAAGAGGGTAT
SEQ2605	TGTTTATTTTGTCCCAATTAATGACCGCCTTTATAAGGGAATAAATGGTAAAGAGGGTAT TGTTTATTTTGTCCCAATTAATGACCGCCTTTATAAGGGAATAAATGGTAAAGAGGGTAT
SEQ2606	TGTTTATTTTGTCCCAATTAATGACCGCCTTTATAAGGGAATAAATGGTAAAGAGGGTAT TGTTTATTTTGTCCCAATTAATGACCGCCTTTATAAGGGAATAAATGGTAAAGAGGGTAT
SEQ2607	TGTTTATTTTGTCCCAATTAATGACCGCCTTTATAAGGGAATAAATGGTAAAGAGGGTAT
SEQ2608	TGTTTATTTTGTCCCAATTAATGACCGCCTTTATAAGGGAATAAATGGTAAAGAGGGTAT
SEQ2609	TGTTTATTTTGTCCCAATTAATGACCGCCTTTATAAGGGAATAAATGGTAAAGAGGGTAT
SEQ2601	TACAGAGTCATCAAATAGTCAGGCAAGTATCACTAATGATGCTCTCTTTACTGGAGACCA
SEQ2602	TATAGAGTCATCAAATAGTCAGGCAAGTATCACTAATGATGCTCTCTTTACTGGAGACCA
SEQ2603	TACAGAGTCATCAAATAGTCAGGCAAGTATCACTAATGATGCTCTCTTTACTGGAGACCA
SEQ2604 SEQ2605	TACAGAGTCATCAAATAGTCAGGCAAGTATCACTAATGATGCTCTCTTTACTGGAGACCA
SE05606	TACAGAGTCATCAAATAGTCAGGCAAGTATCACTAATGATGCTCTCTTTACTGGAGACCA
SEQ2606 SEQ2607	TACAGAGTCATCAAATAGTCAGGCAAGTATCACTAATGATGCTCTCTTTACTGGAGACCA
SEQ2608	TACAGAGTCATCAAATAGTCAGGCAAGTATCACTAATGATGCTCTCTTTACTGGAGACCA
SEQ2609	TACAGAGTCATCAAATAGTCAGGCAAGTATCACTAATGATGCTCTCTTTACTGGAGACCA TACAGAGTCATCAAATAGTCAGGCAAGTATCACTAATGATGCTCTCTTTACTGGAGACCA
	AJJ#BRODITATIOTOTOTOTOTOTOTOTOTOTOTOTOTOTOTOTOTO
SEQ2601	TTTTCATCCCAATAATATTGGCTATCAAATCATGTCTAACGCCGTTATGGAGAAAATAAA
SEQ2602	TTTTCATCCCAATAATATTGGCTATCAAATCATGTCTAACGCCGTTATGGAGAAAATAAA
SEQ2603 '	TTTTCATCCCAATAATATTGGCTATCAAATCATGTCTAACGCCGTTATGGAGAAAATAAA

#### Table 26: Comparative Sequences relating to SAG0503 (lipase/acylhydolase)

SECTION	TTTTCATCCCAATAATATTGGCTATCAAATCATGTCTAACGCCGTTATGGAGAAAATAAA
SEQ2605	TTTTCATCCCAATAATATTGGCTATCAAATCATGTCTAACGCCGTTATGGAGAAAATAAA
SEQ2606	TTTTCATCCCAATAATATTGGCTATCAAATCATGTCTAACGCCGTTATGGAGAAAATAAA
SEQ2607	TTTTCATCCCAATAATATTGGCTATCAAATCATGTCTAACGCCGTTATGGAGAAAATAAA
SEQ2608	TTTTCATCCCAATAATATTGGCTATCAAATCATGTCTAACGCCGTTATGGAGAAAATAAA
SEQ2609	TTTTCATCCCAATAATATTGGCTATCAAATCATGTCTAACGCCGTTATGGAGAAAATAAA
SEQ2601	TGAAACAAGAAAAAACTGGCCGAACCCAGCTTTCTTGTACAAAG
SEQ2602	TGAAACAAGAAAAAACTGGCCGAACCCAGCTTTCTTGTACAAAGTGGTCC
SEQ2603	TGAAACAAGAAAAACTGGCCGAACCCAGCTTTCTTGTACAA
SEQ2604	TGAAACAAGAAAAACTGGCCGAACCCAGCTTTCTTGTACAAA
SEQ2605	TGAAACAAGAAAAACTGGCCGAACCCAGCTTTCTTGTACAA
SEQ2606	TGAAACAAGAAAAACTGGCCGAACCCAGCTTTCTTGTACAAA
SEQ2607	TGAAACAAGAAAAACTGGCCGAACCCAGCTTTCTTGTACAAA
SEQ2608	TGAAACAAGAAAAACTGGCCGAACCCAGCTTTCTTGTACAAAGTGG
SEQ2609	TGAAACAAGAAAAACTGGCCGAACCCAGCTTTCTTGTACAAATABCMARATVSTNCSRA
SEQ2601	~~~~~~~~~
SEQ2602	
SEQ2603	
SEQ2604	
SEQ2605	
SEQ2606	
SEQ2607	
SEQ2608	
SEQ2609	NGTSAGASACYHYDAS

#### >SEQ ID NO 2650:103 090 frame: 2

IFSLIIPKSNPKLTKKDFLTKKVIPLNYVALGDSLTEGVGDTTSQGGFVP
LLSESLHNRYSYQVTSVNYGVSGNTSQQILKRMTTDPQIEKDLEKADLLTLTVGGNDVLA
VIRKELSHLSLNSFEKPAEAYKERLKEILAKARQDNPKLPIYVLGIYNPFYLNFPQLTKM
QTVIDNWNKATKEVVDASENVYFVPINDRLYKGINGKEGITESSNSQASITNDALFTGDH
FHPNNIGYQIMSNAVMEKINETRKNWP

#### >SEQ ID NO 2651:103 H36B frame: 2

IFSLIIPKSNPKLTKKDFLTKKVIPLNYVALGDSLTEGVGDTTSQGGFVPLLS
ESLHNRYSYQVTSVNYGVSGNTSQQILKRMTTDPQIEKDLEKADLLTLTVGGNDVLAVIR
KELSHLSLNSFEKPAEAYKERLKEILAKARQDNPKLPIYVLGIYNPFYLNFPQLTKMQTV
IDNWNKATKEVVDASENVYFVPINDRLYKGINGKEGIIESSNSQASITNDALFTGDHFHP
NNIGYQIMSNAVMEKINETRKNWP

#### >SEQ ID NO 2652:103\_18RS21 frame: 3

IFSLIIPKSNPKLTKKDFLTKKVIPLNYVALGDSLTEGVGDTTSQGGFVPLLS ESLHNRYSYQVTSVNYGVSGNTSQQILKRMTTDPQIEKDLEKADLLTLTVGGNDVLAVIR KELSHLSLNSFEKPAEAYKERLKEILAKARQDNPKLPIYVLGIYNPFYLNFPQLTKMQTV IDNWNKATKEVVDASENVYFVPINDRLYKGINGKEGITESSNSQASITNDALFTGDHFHP NNIGYQIMSNAVMEKINETRKNWP

#### >SEQ ID NO 2653:103 COH1 frame: 3

IFSLIIPKSNPKLTKKDFLTKKVIPLNYVALGDSLTEGVGDTTSQGGFVPL LSESLHNRYSYQVTSVNYGVSGNTSQQILKRMTTDPQIEKDLEKADLLTLTVGGNDVLAV IRKELSHLSLNSFEKPAEAYKERLKEILAKARQDNPKLPIYVLGIYNPFYLNFPQLTKMQ TVIDNWNKATKEVVDASENVYFVPINDRLYKGINGKEGITESSNSQASITNDALFTGDHF HPNNIGYQIMSNAVMEKINETRKNWP

#### >SEQ ID NO 2654:103\_CJB110 frame: 3

IFSLIIPKSNPKLTKKDFLTKKVIPLNYVALGDSLTEGVGDTTSQGGFVPLLS ESLHNRYSYQVTSVNYGVSGNTSQQILKRMTTDPQIEKDLEKADLLTLTVGGNDVLAVIR KELSHLSLNSFEKPAEAYKERLKEILAKARQDNPKLPIYVLGIYNPFYLNFPQLTKMQTV IDNWNKATKEVVDASENVYFVPINDRLYKGINGKEGITESSNSQASITNDALFTGDHFHP NNIGYQIMSNAVMEKINETRKNWP

#### >SEQ ID NO 2655:103\_1169NT frame: 3

IFSLIIPKSNPKLTKKDFLTKKVIPLNYVALGDSLTEGVGDTTSQGGFVPLLS
ESLHNRYSYQVTSVNYGVSGNTSQQILKRMTTDPQIEKDLEKADLLTLTVGGNDVLAVIR
KELSHLSLNSFEKPAEAYKERLKEILAKARQDNPKLPIYVLGIYNPFYLNFPQLTKMQTV
IDNWNKATKEVVDASENVYFVPINDRLYKGINGKEGITESSNSQASITNDALFTGDHFHP
NNIGYQIMSNAVMEKINETRKNWP

### Table 26: Comparative Sequences relating to SAG0503 (lipase/acylhydolase)

### >5 ID NO 2656:103 JM9130013 frame: 3

IFSLIIPKSNPKLTKKDFLTKKVIPLNYVALGDSLTEGVGDTTSQGGFVPLLS
ESLHNRYSYQVTSVNYGVSGNTSQQILKRMTTDPQIEKDLEKADLLTLTVGGNDVLAVIR
KELSHLSLNSFEKPAEAYKERLKEILAKARQDNPKLPIYVLGIYNPFYLNFPQLTKMQTV
IDNWNKATKEVVDASENVYFVPINDRLYKGINGKEGITESSNSQASITNDALFTGDHFHP
NNIGYQIMSNAVMEKINETRKNWP

>SEQ ID NO 2657:103 2603 frame: 1

IFSLIIPKSNPKLTKKDFLTKKVIPLNYVALGDSLTEGVGDTTSQGGFVPLL
SESLHNRYSYQVTSVNYGVSGNTSQQILKRMTTDPQIEKDLEKADLLTLTVGGNDVLAVI
RKELSHLSLNSFEKPAEAYKERLKEILAKARQDNPKLPIYVLGIYNPFYLNFPQLTKMQT
VIDNWNKATKEVVDASENVYFVPINDRLYKGINGKEGITESSNSQASITNDALFTGDHFH
PNNIGYQIMSNAVMEKINETRKNWP

### >SEQ ID NO 2658:103\_M781 frame: 3

IFSLIIPKSNPKLTKKDFLTKKVIPLNYVALGDSLTEGVGDTTSQGGFVPL
LSESLHNRYSYQVTSVNYGVSGNTSQQILKRMTTDPQIEKDLEKADLLTLTVGGNDVLAV
IRKELSHLSLNSFEKPAEAYKERLKEILAKARQDNPKLPIYVLGIYNPFYLNFPQLTKMQ
TVIDNWNKATKEVVDASENVYFVPINDRLYKGINGKEGITESSNSQASITNDALFTGDHF
HPNNIGYQIMSNAVMEKINETRKNWP

SEQ2650	IFSLIIPKSNPKLTKKDFLTKKVIPLNYVALGDSLTEGVGDTTSQGGFVPLLSESLHNRY
SEQ2651	IFSLIIPKSNPKLTKKDFLTKKVIPLNYVALGDSLTEGVGDTTSQGGFVPLLSESLHNRY
SEQ2652	IFSLIIPKSNPKLTKKDFLTKKVIPLNYVALGDSLTEGVGDTTSQGGFVPLLSESLHNRY
SEQ2 <u>6</u> 53	IFSLIIPKSNPKLTKKDFLTKKVIPLNYVALGDSLTEGVGDTTSQGGFVPLLSESLHNRY
SEQ2654	IFSLIIPKSNPKLTKKDFLTKKVIPLNYVALGDSLTEGVGDTTSQGGFVPLLSESLHNRY
SEQ2655	IFSLIIPKSNPKLTKKDFLTKKVIPLNYVALGDSLTEGVGDTTSQGGFVPLLSESLHNRY
SEQ2656	IFSLIIPKSNPKLTKKDFLTKKVIPLNYVALGDSLTEGVGDTTSQGGFVPLLSESLHNRY
SEQ2657	IFSLII PKSNPKLTKKDFLTKKVI PLNYVALGDSLTEGVGDTTSQGGFVPLLSESLHNRY
SEQ2658	IFSLIIPKSNPKLTKKDFLTKKVIPLNYVALGDSLTEGVGDTTSQGGFVPLLSESLHNRY
SEQ2650	SYQVTSVNYGVSGNTSQQILKRMTTDPQIEKDLEKADLLTLTVGGNDVLAVIRKELSHLS
SEQ2651	SYQVTSVNYGVSGNTSQQILKRMTTDPQIEKDLEKADLLTLTVGGNDVLAVIRKELSHLS
SEQ2652	SYQVTSVNYGVSGNTSQQILKRMTTDPQIEKDLEKADLLTLTVGGNDVLAVIRKELSHLS
SEQ2653	SYQVTSVNYGVSGNTSQQILKRMTTDPQIEKDLEKADLLTLTVGGNDVLAVIRKELSHLS
SEQ2654	SYQVTSVNYGVSGNTSQQILKRMTTDPQIEKDLEKADLLTLTVGGNDVLAVIRKELSHLS
SEQ2655	SYQVTSVNYGVSGNTSQQILKRMTTDPQIEKDLEKADLLTLTVGGNDVLAVIRKELSHLS
SEQ2656	SYQVTSVNYGVSGNTSQQILKRMTTDPQIEKDLEKADLLTLTVGGNDVLAVIRKELSHLS
SEQ2657	SYQVTSVNYGVSGNTSQQILKRMTTDPQIEKDLEKADLLTLTVGGNDVLAVIRKELSHLS
SEQ2658	
22220	SYQVTSVNYGVSGNTSQQILKRMTTDPQIEKDLEKADLLTLTVGGNDVLAVIRKELSHLS
SEQ2650	I NCERVARA VVERY VETT AVERA DODUNION DE LUCIO CENTRA DE LA CONTRA DEL CONTRA DE LA CONTRA DEL CONTRA DE LA CONTRA DEL CONTRA DE LA CONTRA DEL CONTRA DE LA CONTRA DEL CONTRA DE LA CONTRA D
SEQ2651	LNSFEKPAEAYKERLKEILAKARQDNPKLPIYVLGIYNPFYLNFPQLTKMQTVIDNWNKA
SEQ2652	LNSFEKPAEAYKERLKEILAKARQDNPKLPIYVLGIYNPFYLNFPQLTKMQTVIDNWNKA
_	LNSFEKPAEAYKERLKEILAKARQDNPKLPIYVLGIYNPFYLNFPQLTKMQTVIDNWNKA
SEQ2653	LNSFEKPAEAYKERLKEILAKARQDNPKLPIYVLGIYNPFYLNFPQLTKMQTVIDNWNKA
SEQ2654	LNSFEKPAEAYKERLKEILAKARQDNPKLPIYVLGIYNPFYLNFPQLTKMQTVIDNWNKA
SEQ2655	LNSFEKPAEAYKERLKEILAKARQDNPKLPIYVLGIYNPFYLNFPQLTKMQTVIDNWNKA
SEQ2656	LNSFEKPAEAYKERLKEILAKARQDNPKLPIYVLGIYNPFYLNFPQLTKMQTVIDNWNKA
SEQ2657	LNSFEKPAEAYKERLKEILAKARQDNPKLPIYVLGIYNPFYLNFPQLTKMQTVIDNWNKA
SEQ2658	LNSFEKPAEAYKERLKEILAKARQDNPKLPIYVLGIYNPFYLNFPQLTKMQTVIDNWNKA
SEQ2650	TKEVVDASENVYFVPINDRLYKGINGKEGITESSNSQASITNDALFTGDHFHPNNIGYQI
SEQ2651	TKEVVDASENVYFVPINDRLYKGINGKEGIIESSNSQASITNDALFTGDHFHPNNIGYQI
SEQ2652	TKEVVDASENVYFVPINDRLYKGINGKEGITESSNSQASITNDALFTGDHFHPNNIGYQI
SEQ2653	TKEVVDASENVYFVPINDRLYKGINGKEGITESSNSQASITNDALFTGDHFHPNNIGYQI
SEQ2654	TKEVVDASENVYFVPINDRLYKGINGKEGITESSNSQASITNDALFTGDHFHPNNIGYQI
SEQ2655	TKEVVDASENVYFVPINDRLYKGINGKEGITESSNSQASITNDALFTGDHFHPNNIGYQI
SEQ2656	TKEVVDASENVYFVPINDRLYKGINGKEGITESSNSQASITNDALFTGDHFHPNNIGYQI
SEQ2657	TKEVVDASENVYFVPINDRLYKGINGKEGITESSNSQASITNDALFTGDHFHPNNIGYQI
SEQ2658	TKEVVDASENVYFVPINDRLYKGINGKEGITESSNSQASITNDALFTGDHFHPNNIGYQI
SEQ2650	MSNAVMEKINETRKNWP
SEQ2651	MSNAVMEKINETRKNWP
SEQ2652	MSNAVMEKINETRKNWP
SEQ2653	MSNAVMEKINETRKNWP
SEQ2654	MSNAVMEKINETRKNWP
SEQ2655	MSNAVMEKINETRKNWP
SEQ2656	
SEQ2657	MSNAVMEKINETRKNWP
SEQ2658	MSNAVMEKINETRKNWP
2H2E030	MSNAVMEKINETRKNWP

## Tabl 27: C mparative Sequences relating t SAG1473 (cell wall surface anchor family protein)

SEQ ID NO. 2701: SAG1473 FROM THE 1169NT1 GBS TYPE V STRAIN (REVERSE COMPLEMENT)

GATACAAGTGATAAGAATACTGACACGAGTGTCGTGACTACGACCTTATCTGAGGAGAAAAGATCAGATGA ACTAGACCAGTCTAGTACTGGTTCTTCTTCTGAAAATGAATCGAGTTCATCAAGTGAACCAGAAACCAAATC CGTCAACTAATCCACCTACAACAGAACCATCGCAACCCTCACCTAGTGAAGAAAAAGCCTGATGGTAGA ACGAAGACAGAAATTGGCAATAATAAGGATATTTCTAGTGGAACAAAAGTATTAATTTCAGAAGATAGTAT TAAGAATTTTAGTAAAGCAAGTAGTGATCAAGAAGAAGAAGTAGTATCATCATCATCAAAAAGCAA GTGATGGGAAAAAAAGGCCAACAGTAAGCCTAAAAAAGGAA

SEQ ID NO. 2702: SAG1473 FROM THE 18RS21 GBS TYPE II STRAIN
GATACAAGTGATAAGAATACTGACACGAGTGTCGTGACTACGACCTTATCTGAGGAGAAAAGATCAGATGA
ACTAGACCAGTCTAGTACTGGTTCTTCTTCTGAAAATGAATCGAGTTCATCAAGTGAACCAGAAACAAATC
CGTCAACTAATCCACCTACAACAGAACCATCGCAACCCTCACCTAGTGAAGAAAAAGCCTGATGGTAGA
ACGAAGACAGAAATTGGCAATAATAAGGATATTTCTAGTGGAACAAAAGTATTAATTTCAGAAGATAGTAT
TAAGAATTTTAGTAAAGCAAGTAGTGATCAAGAAGAAGTGGATCGCGATGAATCATCATCTTCAAAAAGCAA
ATGATGGGAAAAAAAGGCCACAGTAAGCCTAAAAAGGAA

SEQ ID NO. 2703: SAG1473 FROM THE 2603 V/R GBS TYPE V STRAIN
GATACAAGTGATAAGAATACTGACACGAGTGTCGTGACTACGACCTTATCTGAGGAGAAAAGATCAGATGA
ACTAGACCAGTCTAGTACTGGTTCTTCTTCTGAAAATGAATCGAGTTCATCAAGTGAACCAGAAACAAATC
CGTCAACTAATCCACCTACAACAGAACCATCGCAACCCTCACCTAGTGAAGAGAACAAGCCTGATGGTAGA
ACGAAGACAGAAATTGGCAATAATAAGGATATTTCTAGTGGAACAAAAGTATTAATTTCAGAAGATAGTAT
TAAGAATTTTAGTAAAGCAAGTAGTGATCAAGAAGAAGTGGATCGCGATGAATCATCATCTTCAAAAAGCAA
ATGATGGGAAAAAAAGGCCACAGTAAGCCTAAAAAGGAA

SEQ ID NO. 2704: SAG1473 FROM THE 090 GBS TYPE IA STRAIN
GACCAGTCTAGTACTGGTTCTTCTTCTGAAAATGAATCGAGTTCATCAAGTGAACCAGAAACAAATCCGTC
AACTAATCCACCTACAACAGAACCATCGCAACCCTCACCTAGTGAAGAACAAGCCTGATGGTAGAACGA
AGACAGAAATTGGCAATAATAAGGATATTTCTAGTGGAACAAAAGTATTAATTTCAGAAGATAGTATTAAG
AATTTTAGTAAAGCAAGTAGTGATCAAGAAGAAGTGGATCGCGATGAATCATCATCTTCAAAAGCAAATGA
TGGGAAAAAAGGCCACAGTAAGCCTAAAAAGGAA

SEQ ID NO. 2705: SAG1473 FROM THE A909 GBS TYPE Ia STRAIN , GATACAAGTGATAAGAATACTGACACGAGTGTCGTGACTACGACCTTATCTGAGGAGAAAAGATTAGATGA ACTAGACCAGTCTAGTACTGGTTCTTCTTCTGAAAATGAATCGAGTTCATCAAGTGAACCAGAAACCAAATC CCTCAACTAATCCACCTACAACAGAACCATCGCAACCCTCACCTAGTGAAGAAACAAGCCTGATGGTAGC ACGAAGACAGAAATTGGCAATAATAAGGATATTTCTAGTGGAACAAAAGTATTAATTTCAGAAGATAGTAT TAAGAATTTTAGTAAAGCAAGTAGTGATCAAGAAGAAGAAGTAGTATCATCATCATCAAAAAGCAA ATGATGAGAAAAAAAGGCCCACAGTAAGCCTAAAAAGGAA

SEQ ID NO. 2706: SAG1473 FROM THE CJB110 GBS NONTYPEABLE STRAIN GATACAAGTGATAAGAATACTGACACGAGTGTCGTGACTACGACCTTATCTGAGGAGAAAAGATCAGATGA ACTAGACCAGTCTAGTTACTGGTTCTTCTTCTGAAAATGAATCGAGTTCATCAAGTGAACCAGAAACAAATC CGTCAACTAATCCACCTACAACAGAACCATCGCAACCCTCACCTAGTGAAGAAAAAAGCCTGATGGTAGA ACGAAGACAGAAATTGGCAATAATAAGGATATTTCTAGTGGAACAAAAGTATTAATTTCAGAAGATAGTAT TAAGAATTTTAGTAAAAGCAAGTAGTGATCAAGAAGAAGAAAAGCAA ATGATGGGAAAAAAAGCCACAGTAAGCCTAAAAAGGAA

## Table 27: Comparative Sequences relating t SAG14/3 (cell wall surface anch r family protein)

### SEQ ID NO. 2707: SAG1473 FROM THE COH1 GBS TYPE III STRAIN (REVERSE COMPLEMENT)

GATACAAGTGATAAGAATACTGACACGAGTGTCGTGACTACGACCTTATCTGAGGAGAAAAGATCAGATGA
ACTAGACCAGTCTAGTACTGGTTCTTCTTCTGAAAATGAATCAAGTTCATCAAGTGAACCAGAAACAAATC
CCTCAACTAATCCACCTACAACAGAACCATCGCAACCCTCACCTAGTGAAGAGAACAAGCCTGATGGGAGC
ACGAAGACAGAAATTGGCAATAATAAGGATATTTCTAGTGGAACAAAAGTATTAATTTCAGAAGATAGTAT
TAAGAATTTTAGTAAAGCAAGTAGTGATCAAGAAGAAGTGGAACGCGATGAATCATCATCTTCAAAAGCAA
ATGATGAGAAAAAAAGGCCACAGTAAGCCTAAAAAGGAA

### SEQ ID NO. 2708: SAG1473 FROM THE H36b GBS TYPE Ib STRAIN

GATACAAGTGATAAGAATACTGACACGAGTGTCGTGACTACGACCTTATCTGAGGAGAAAAGATTAGATGA
ACTAGACCAGTCTAGTACTGGTTCTTCTTCTGAAAATGAATCGAGTTCATCAAGTGAACCAGAAACAAATC
CCTCAACTAATCCACCTACAACAGAACCATCGCAACCCTCACCTAGTGAAGAACAAGCCTGATGGTAGC
ACGAAGACAGAAATTGGCAATAATAAGGATATTTCTAGTGGAACAAAAGTATTAATTTCAGAAGATAGTAT
TAAGAATTTTAGTAAAGCAAGTAGTGATCAAGAArAAGGATCGCGATGAATCATCATCATCAAAAAGCAA
ATGATGAGAAAAAAAGGCCACAGTAAGCCTAAAAAGGAA

### SEQ ID NO. 2709: SAG1473 FROM THE JM910013 GBS TYPE VIII STRAIN

GATACAAGTGATAAGAATACTGACACGAGTGTCGTGACTACGACCTTATCTGAGGAGAAAAGATTAGATGA
ACTAGACCAGTCTAGTACTGGTTCTTCTTCTGAAAATGAATCGAGTTCATCAAGTGAACCAGAAACAAATC
CCTCAACTAATCCACCTACAACAGAACCATCGCAACCCTCACCTAGTGAAGAGAACAAGCCTGATGGTAGC
ACGAAGACAGAAATTGGCAATAATAAGGATATTTCTAGTGGAACAAAAGTATTAATTTCAGAAGATAGTAT
TAAGAATTTTAGTAAAAGCAAGTAGTGATCAAGAAGAAGTGGATCGTCATCATCATCAAAAAGCAA
ATGATGAGAAAAAAAGGCCACAGTAAGCCTAAAAAGGAA

### SEQ ID NO. 2710: SAG1473 FROM THE M732 GBS TYPE III STRAIN

GATACAAGTGATAAGAATACTGACACGAGTGTCGTGACTACGACCTTATCTGAGGAGAAAAGATCAGATGA ACTAGACCAGTCTAGTACTGGTTCTTCTTCTGAAAATGAATCAAGTTCATCAAGTGAACCAGAAACAAATC CCTCAACTAATCCACCTACAACAGAACCATCGCAACCCTCACCTAGTGAAGAGAACAAGCCTGATGGGAGC ACGAAGACAGAAATTGGCAATAATAAGGATATTTCTAGTGGAACAAAAGTATTAATTTCAGAAGATAGTAT TAAGAATTTTAGTAAAGCAAGTAGTGATCAAGAAGAAGTGGAACGCGATGAATCATCATCTTCAAAAAGCAA ATGATGAGAAAAAAGGCCACAGTAAGCCTAAAAAGGAA

### SEQ ID NO. 2711: SAG1473 FROM THE M781 GBS TYPE III STRAIN

GATACAAGTGATAAGAATACTGACACGAGTGTCGTGACTACGACCTTATCTGAGGAGAAAAGATCAGATGA ACTAGACCAGTCTAGTACTGGTTCTTCTTCTGAAAATGAATCAAGTTCATCAAGTGAACCAGAAACAAATC CCTCAACTAATCCACCTACAACAGAACCATCGCAACCCTCACCTAGTGAAGAACAAGCCTGATGGGAGC ACGAAGACAGAAATTGGCAATAATAAGGATATTTCTAGTGGAACAAAAGTATTAATTTCAGAAGATAGTAT TAAGAATTTTAGTAAAGCAAGTAGTGATCAAGAAGAAGTGGATCGCGATGAATCATCATCTTCAAAAGCAA ATGATGAGAAAAAAAGGCCACAGTAAGCCTAAAAAGGAA

SEQ2701	atacaagtgataagaatactgacacgagtgtcgtgactacgaccttatctgaggagaaa
SEQ2702	atacaagtgataagaatactgacacgagtgtcgtgactacgaccttatctgaggagaaa
SEQ2703	ATACAAGTGATAAGAATACTGACACGAGTGTCGTGACTACGACCTTATCTGAGGAGAAA
SEQ2704	
SEQ2705	ATACAAGTGATAAGAATACTGACACGAGTGTCGTGACTACGACCTTATCTGAGGAGAAA
SEQ2706	ATACAAGTGATAAGAATACTGACACGAGTGTCGTGACTACGACCTTATCTGAGGAGAAA
SEQ2707	ATACAAGTGATAAGAATACTGACACGAGTGTCGTGACTACGACCTTATCTGAGGAGAAA
SEQ2709	ATACAAGTGATAAGAATACTGACACGAGTGTCGTGACTACGACCTTATCTGAGGAGAAA
SEQ2710	ATACAAGTGATAAGAATACTGACACGAGTGTCGTGACTACGACCTTATCTGAGGAGAAA
SEQ2711	ATACAAGTGATAAGAATACTGACACGAGTGTCGTGACTACGACCTTATCTGAGGAGAAA



# Table 27: C mparative Sequences relating to SAG1473 (cell wall surface anchor family protein)

SEQ2701	GATCAGATGAACTAĞACCAGTCTAGTACTGGTTCTTCTTCTGAAAATGAATCGAGTTCA
SEQ2702	GATCAGATGAACTAGACCAGTCTAGTACTGGTTCTTCTTCTGAAAATGAATCGAGTTCA
SEQ2703	GATCAGATGAACTAGACCAGTCTAGTACTGGTTCTTCTTCTGAAAATGAATCGAGTTCA
SEQ2704	
GACCAGTCTAGTACT	GGTTCTTCTGAAAATGAATCGAGTTCA
SEQ2705	GATTAGATGAACTAGACCAGTCTAGTACTGGTTCTTCTTCTGAAAATGAATCGAGTTCA
SEQ2706	GATCAGATGAACTAGACCAGTCTAGTACTGGTTCTTCTTCTGAAAATGAATCGAGTTCA
SEQ2707	GATCAGATGAACTAGACCAGTCTAGTACTGGTTCTTCTTCTGAAAATGAATCAAGTTCA
SEQ2709	GATTAGATGAACTAGACCAGTCTAGTACTGGTTCTTCTTCTGAAAATGAATCGAGTTCA
SEQ2710	GATCAGATGAACTAGACCAGTCTAGTACTGGTTCTTCTTCTGAAAATGAATCAAGTTCA
SEQ2711	GATCAGATGAACTAGACCAGTCTAGTACTGGTTCTTCTTGTAAAATGAATCAAGTTCA
SEQ2701	TCAAGTGAACCAGAAACAAATCCGTCAACTAATCCACCTACAACAGAACCATCGCAACCC
SEQ2702	TCAAGTGAACCAGAAACAAATCCGTCAACTAATCCACCTACAACAGAACCATCGCAACCC
SEQ2703	TCAAGTGAACCAGAAACAAATCCGTCAACTAATCCACCTACAACAGAACCATCGCAACCC
SEQ2704	TCAAGTGAACCAGAAACAAATCCGTCAACTAATCCACCTACAACAGAACCATCGCAACCC
SEQ2705	TCAAGTGAACCAGAAACAAATCCCTCAACTAATCCACCTACAACAGAACCATCGCAACCC
SEQ2706	TCAAGTGAACCAGAAACAAATCCGTCAACTAATCCACCTACAACAGAACCATCGCAACCC
SEQ2707	TCAAGTGAACCAGAAACAAATCCCTCAACTAATCCACCTACAACAGAACCATCGCAACCC
SEQ2709	TCAAGTGAACCAGAAACAAATCCCTCAACTAATCCACCTACAACAGAACCATCGCAACCC
SEQ2710	TCAAGTGAACCAGAAACAAATCCCTCAACTAATCCACCTACAACAGAACCATCGCAACCC
SEQ2711	TCAAGTGAACCAGAAACAAATCCCTCAACTAATCCACCTACAACAGAACCATCGCAACCC
SEQ2701	TCACCTAGTGAAGAGAACAAGCCTGATGGTAGAACGAAGACAGAAATTGGCAATAATAAG
SEQ2702	TCACCTAGTGAAGAGAACAAGCCTGATGGTAGAACGAAGACAGAAATTGGCAATAATAAG
SEQ2703	TCACCTAGTGAAGAGAACAAGCCTGATGGTAGAACGAAGACAGAAATTGGCAATAATAAG
SEQ2704	TCACCTAGTGAAGAGAACAAGCCTGATGGTAGAACGAAGACAGAAATTGGCAATAATAAG
SEQ2705	TCACCTAGTGAAGAGAACAAGCCTGATGGTAGCACGAAGACAGAAATTGGCAATAATAAG
SEQ2706	TCACCTAGTGAAGAGAACAAGCCTGATGGTAGAACGAAGACAGAAATTGGCAATAATAAG
SEQ2707	TCACCTAGTGAAGAGAACAAGCCTGATGGGAGCACGAAGACAGAAATTGGCAATAATAAG
SEQ2709	TCACCTAGTGAAGAGAACAAGCCTGATGGTAGCACGAAGACAGAAATTGGCAATAATAAG
SEQ2710	TCACCTAGTGAAGAGAACAAGCCTGATGGGAGCACGAAGACAGAAATTGGCAATAATAAG
SEQ2711	TCACCTAGTGAAGAGAACAAGCCTGATGGGAGCACGAAGACAGAAATTGGCAATAATAAG
SEQ2701	GATATTTCTAGTGGAACAAAAGTATTAATTTCAGAAGATAGTATTAAGAATTTTAGTAAA
SEQ2702	GATATTTCTAGTGGAACAAAAGTATTAATTTCAGAAGATAGTATTAAGAATTTTAGTAAA
SEQ2703	GATATTTCTAGTGGAACAAAAGTATTAATTTCAGAAGATAGTATTAAGAATTTTAGTAAA
SEQ2704	GATATTTCTAGTGGAACAAAAGTATTAATTTCAGAAGATAGTATTAAGAATTTTAGTAAA
SEQ2705	GATATTTCTAGTGGAACAAAAGTATTAATTTCAGAAGATAGTATTAAGAATTTTAGTAAA
SEQ2706	GATATTTCTAGTGGAACAAAAGTATTAATTTCAGAAGATAGTATTAAGAATTTTAGTAAA
SEQ2707	GATATTTCTAGTGGAACAAAAGTATTAATTTCAGAAGATAGTATTAAGAATTTTAGTAAA
SEQ2709	GATATTTCTAGTGGAACAAAAGTATTAATTTCAGAAGATAGTATTAAGAATTTTAGTAAA
SEQ2710	GATATTTCTAGTGGAACAAAAGTATTAATTTCAGAAGATAGTATTAAGAATTTTAGTAAA
SEQ2711	GATATTTCTAGTGGAACAAAAGTATTAATTTCAGAAGATAGTATTAAGAATTTTAGTAAA
SEQ2701	GCAAGTAGTGATCAAGAAGAAGTGGATCGCGATGAATCATCTTCAAAAGCAAGTGAT
SEQ2702	GCAAGTAGTGATCAAGAAGAAGTGGATCGCGATGAATCATCATCTTCAAAAGCAAATGAT
SEQ2703	GCAAGTAGTGATCAAGAAGAAGTGGATCGCGATGAATCATCTTCAAAAGCAAATGAT
SEQ2704	GCAAGTAGTGATCAAGAAGAAGTGGATCGCGATGAATCATCTTCAAAAGCAAATGAT
SEQ2705	GCAAGTAGTGATCAAGAAGAAGTGGATCGCGATGAATCATCTTCAAAAGCAAATGAT
SEQ2706	GCAAGTAGTGATCAAGAAGAAGTGGATCGCGATGAATCATCTTCAAAAGCAAATGAT
SEQ2707	GCAAGTAGTGATCAAGAAGAAGTGGAACGCGATGAATCATCATCTTCAAAAGCAAATGAT
SEQ2709	GCAAGTAGTGATCAAGAAGAAGTGGATCGCGATGAATCATCATCTTCAAAAGCAAATGAT
SEQ2710	GCAAGTAGTGATCAAGAAGAAGTGGAACGCGATGAATCATCATCTTCAAAAGCAAATGAT
SEQ2711	GCAAGTAGTGATCAAGAAGAAGTGGATCGCGATGAATCATCATCTTCAAAAGCAAATGAT



# Table 27: Comparative Sequences relating t SAG1473 (cell wall surface anchor family protein)

SE02701	GGGAAAAAGGCCACAGTAAGCCTAAAAAGGAA
SEQ2702	GGGAAAAAGGCCACAGTAAGCCTAAAAAGGAA
SEQ2703	GGGAAAAAAGGCCACAGTAAGCCTAAAAAGGAA
SEQ2704	GGGAAAAAGGCCACAGTAAGCCTAAAAAGGAA
SEQ2705	GAGAAAAAAGGCCACAGTAAGCCTAAAAAGGAA
SEQ2706	GGGAAAAAAGGCCACAGTAAGCCTAAAAAGGAA
SEQ2707	GAGAAAAAAGGCCACAGTAAGCCTAAAAAGGAASGATACAAGTGATAAGAATACTGACAC
SEQ2709	GAGAAAAAAGGCCACAGTAAGCCTAAAAAGGAA
SEQ2710	GAGAAAAAGGCCACAGTAAGCCTAAAAAGGAA
SEQ2711	GAGAAAAAAGGCCACAGTAAGCCTAAAAAGGAATABCMARATVSTNCSRATNGTSAGCWA
SEQ2 / II	
SEQ2701	
SEQ2702	
SEQ2702 SEQ2703	
SEQ2703 SEQ2704	
SEQ2705	
SEQ2706	AGTGTCGTGACTACGACCTTATCTGAGGAGAAAAGATTAGATGAACTAGACCAGTCTAG
SEQ2707	AGTGTCGTGACTACCACCTTATCTGAGGAGAAAAGATTAGATGAACTAGACCAGTCTAG
SEQ2709	
SEQ2710	
SEQ2711	TRACANCHRAMYRTN
SEQ2701	
SEQ2702	
SEQ2703	
SEQ2704	
SEQ2705	#
SEQ2706	
SEQ2707	ACTGGTTCTTCTGAAAATGAATCGAGTTCATCAAGTGAACCAGAAACAAATCCCTC
SEQ2709	
SEQ2710	
SEQ2711	~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~
SEQ2701	
SEQ2702	
SEQ2703	
SEQ2704	
SEQ2705	
SEQ2706	
SEQ2707	ACTAATCCACCTACAACAGAACCATCGCAACCCTCACCTAGTGAAGAGAACAAGCCTGA
SEQ2709	
SEQ2710	
SEQ2711	
SEQ2701	
SEQ2701	
SEQ2702 SEQ2703	
SEQ2703 SEQ2704	
<del></del>	
SEQ2705	
SEQ2706	
SEQ2707	GGTAGCACGAAGACAGAAATTGGCAATAATAAGGATATTTCTAGTGGAACAAAAGTATT
SEQ2709	
SEQ2710	
SEQ2711	######################################

# Table 27: C mparative Sequences relating to SAG1473 (cell wall surface anchor family protein)

SEQ2701	
SEQ2702	
SEQ2703	
SEQ2704	, 
SEQ2705	
SEQ2706	
SEQ2707	ATTTCAGAAGATAGTATTAAGAATTTTAGTAAAGCAAGTAGTGATCAAGAARAAGTGGA
SEQ2709	**************************************
SEQ2710	***************************************
SEQ2711	***************************************
SEQ2701	
SEQ2702	***************************************
SEQ2703	
SEQ2704	
SEQ2705	
SEQ2706	
SEQ2707	CGCGATGAATCATCATCTTCAAAAGCAAATGATGAGAAAAAAGGCCACAGTAAGCCTAA
SEQ2709	+
SEQ2710	
SEQ2711	
SEQ2701	=====
SEQ2702	# C C C C C C C C C C C C C C C C C C C
SEQ2703	
SEQ2704	
SEQ2705	
SEQ2706	
SEQ2707	AAGGAA
SEQ2709	
SEQ2710	
SEQ2711	

### >SEQ ID NO 2750:4\_1169NT frame: 1

DTSDKNTDTSVVTTTLSEEKRSDELDQSSTGSSSENESSSSEPETNPSTNPPTTEPSQP SPSEENKPDGRTKTEIGNNKDISSGTKVLISEDSIKNFSKASSDQEEVDRDESSSSKASD GKKGHSKPKKE

### >SEQ ID NO 2751:4\_18RS21 frame: 1

DTSDKNTDTSVVTTTLSEEKRSDELDQSSTGSSSENESSSSSEPETNPSTNPPTTEPSQP SPSEENKPDGRTKTEIGNNKDISSGTKVLISEDSIKNFSKASSDQEEVDRDESSSSKAND GKKGHSKPKKE

### >SEQ ID NO 2752:4\_2603 frame: 1

DTSDKNTDTSVVTTTLSEEKRSDELDQSSTGSSSENESSSSSEPETNPSTNPPTTEPSQP SPSEENKPDGRTKTEIGNNKDISSGTKVLISEDSIKNFSKASSDQEEVDRDESSSSKAND GKKGHSKPKKE

### >SEQ ID NO 2753:4\_090 frame: 1

DQSSTGSSSENESSSSEPETNPSTNPPTTEPSQPSPSEENKPDGRTKTEIGNNKDISSG TKVLISEDSIKNFSKASSDQEEVDRDESSSSKANDGKKGHSKPKKE

### >SEQ ID NO 2754:4\_A909 frame: 1

DTSDKNTDTSVVTTTLSEEKRLDELDQSSTGSSSENESSSSSEPETNPSTNPPTTEPSQP SPSEENKPDGSTKTEIGNNKDISSGTKVLISEDSIKNFSKASSDQEEVDRDESSSKAND EKKGHSKPKKE

### >SEQ ID NO 2755:4\_CJB110 frame: 1

DTSDKNTDTSVVTTTLSEEKRSDELDQSSTGSSSENESSSSSEPETNPSTNPPTTEPSQP SPSEENKPDGRTKTEIGNNKDISSGTKVLISEDSIKNFSKASSDQEEVDRDESSSKAND GKKGHSKPKKE

## Table 27: Comparative Sequences r lating to SAG1473 (cell wall surface anch r family protein)

#### >SEQ ID NO 2756:4 COH1 frame: 1

DTSDKNTDTSVVTTTLSEEKRSDELDQSSTGSSSENESSSSEPETNPSTNPPTTEPSQP SPSEENKPDGSTKTEIGNNKDISSGTKVLISEDSIKNFSKASSDQEEVERDESSSSKAND EKKGHSKPKKE

### >SEQ ID NO 2757:4\_H36B frame: 1

DTSDKNTDTSVVTTTLSEEKRLDELDQSSTGSSSENESSSSEPETNPSTNPPTTEPSQP SPSEENKPDGSTKTEIGNNKDISSGTKVLISEDSIKNFSKASSDQEXVDRDESSSSKAND EKKGHSKPKKE

#### >SEQ ID NO 2758:4 JM9130013 frame: 1

DTSDKNTDTSVVTTTLSEEKRLDELDQSSTGSSSENESSSSSEPETNPSTNPPTTEPSQP SPSEENKPDGSTKTEIGNNKDISSGTKVLISEDSIKNFSKASSDQEEVDRDESSSSKAND EKKGHSKPKKE

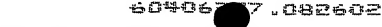
### >SEQ ID NO 2759:4 M732 frame: 1

DTSDKNTDTSVVTTTLSEEKRSDELDQSSTGSSSENESSSSSEPETNPSTNPPTTEPSQP SPSEENKPDGSTKTEIGNNKDISSGTKVLISEDSIKNFSKASSDQEEVERDESSSSKAND EKKGHSKPKKE

### >SEQ ID NO 2760:4\_M781 frame: 1

DTSDKNTDTSVVTTTLSEEKRSDELDQSSTGSSSENESSSSSEPETNPSTNPPTTEPSQP SPSEENKPDGSTKTEIGNNKDISSGTKVLISEDSIKNFSKASSDQEEVDRDESSSSKAND EKKGHSKPKKE

SEQ2750	TSDKNTDTSVVTTTLSEEKRSDELDQSSTGSSSENESSSSSEPETNPSTNPPTTEPSQP
SEQ2751	TSDKNTDTSVVTTTLSEEKRSDELDQSSTGSSSENESSSSSEPETNPSTNPPTTEPSQP
SEQ2752	TSDKNTDTSVVTTTLSEEKRSDELDQSSTGSSSENESSSSSEPETNPSTNPPTTEPSQP
SEQ2753	
SEQ2754	TSDENTEDTS SHIPTED SEEVEL DEL DOSSESSENESSSSEPETNESTNERTEPS QP
SEQ2755	TSDKNTDTSVVTTTLSEEKRLDELDQSSTGSSSENESSSSSEPETNPSTNPPTTEPSQP
SEQ2756	TSDKNTDTSVVTTTLSEEKRSDELDQSSTGSSSENESSSSSEPETNPSTNPPTTEPSQP
SEQ2757	TSDKNTDTSVVTTTLSEEKRSDELDQSSTGSSSENESSSSSEPETNPSTNPPTTEPSQP
SEQ2758	TSDKNTDTSVVTTTLSEEKRLDELDQSSTGSSSENESSSSSEPETNPSTNPPTTEPSQP
SEQ2759	TSDKNTDTSVVTTTLSEEKRLDELDQSSTGSSSENESSSSSEPETNPSTNPPTTEPSQP
SEQ2760	TSDKNTDTSVVTTTLSEEKRSDELDQSSTGSSSENESSSSSEPETNPSTNPPTTEPSQP
3EQ2 / 60	TSDKNTDTSVVTTTLSEEKRSDELDQSSTGSSSENESSSSSEPETNPSTNPPTTEPSQP
SEQ2750	
SEQ2751	SPSEENKPDGRTKTEIGNNKDISSGTKVLISEDSIKNFSKASSDQEEVDRDESSSSKASD
SEQ2752	SPSEENKPDGRTKTEIGNNKDISSGTKVLISEDSIKNFSKASSDQEEVDRDESSSSKAND
SEQ2753	SPSEENKPDGRTKTEIGNNKDISSGTKVLISEDSIKNFSKASSDQEEVDRDESSSSKAND
SEQ2754	SPSEENKPDGRTKTEIGNNKDISSGTKVLISEDSIKNFSKASSDQEEVDRDESSSSKAND
SEQ2755	SPSEENKPDGSTKTEIGNNKDISSGTKVLISEDSIKNFSKASSDQEEVDRDESSSSKAND
SEQ2756	SPSEENKPDGRTKTEIGNNKDISSGTKVLISEDSIKNFSKASSDQEEVDRDESSSSKAND
SEQ2757	SPSEENKPDGSTKTEIGNNKDISSGTKVLISEDSIKNFSKASSDQEEVERDESSSSKAND
	SPSEENKPDGSTKTEIGNNKDISSGTKVLISEDSIKNFSKASSDQEXVDRDESSSSKAND
SEQ2758	SPSEENKPDGSTKTEIGNNKDISSGTKVLISEDSIKNFSKASSDQEEVDRDESSSSKAND
SEQ2759	SPSEENKPDGSTKTEIGNNKDISSGTKVLISEDSIKNFSKASSDQEEVERDESSSSKAND
SEQ2760	SPSEENKPDGSTKTEIGNNKDISSGTKVLISEDSIKNFSKASSDQEEVDRDESSSSKAND
SEQ2750	KKGHSKPKKE
SEQ2751	KKGHSKPKKE
SEQ2752	KKGHSKPKKE
SEQ2753	KKGHSKPKKE
SEQ2754	KKGHSKPKKE
SEQ2755	KKGHSKPKKE
SEQ2756	KKGHSKPKKE
SEQ2757	KKGHSKPKKE
SEQ2758	KKGHSKPKKE
SEQ2759	KKGHSKPKKE
SEQ2760	KKGHSKPKKE



# Table 20: C mparative Sequences relating to SAG1552 (conserved hypothetical protein)

SEQ ID NO. 2801: SAG1552 FROM THE 1169NT1 GBS TYPE V STRAIN (REVERSE COMPLEMENT)

TTTGTTGTTAAAGGTGATACTGTACTTCACAAGCCCACCAATAAACCTTTTGTTGTTAAAGGAGTAGACGTTGAGTCT TCCTTAGCAGGTTATCATCACAACGATTTTCCTATTACTCAAAAAACGTATCGTGAGTGGTTCCATTTAATTTCCAAC ATGGGGGCAAATACTGTAAGAGTCAAAGTACCGATGAATGTTGCATTTTACGATGCTTTATATCACCACAACAAAGCA TCAAAGAGGCCACTGTATTTGTTGCAAGGAATACGTATAGATTCTTATCGCAATAATGCTTCTATAACAGCTTTTAAT GATAATTATAGGGGGTATTTAAAACGAGAAGCAAAAGGCGTTGTGGATATTCTCCATGGGCGTAAGCAAGTATGGAAT ACTGATTTTGGTAGCCGTCATTATCATTATGATCTTAGTCCTTGGGTACTTGGTTATGTCGTAGGGGATGATTGGAAT AGTGGTACTGTCGCTTATACTAATCATCAAGAGAAAAAAACGCAATATAAAGGACGTTATTTTAAAACTTCTGCGGCA GCTAATCCATTTGAGGTCATGCTAGCTCAAGTTATGGATGAATTGACACATTATGAGACAGCTAAATATGGTTGGCAA CATTTGATTAGTTTTCAAACTCACCAACAACAGACCCTTTTCGTTATCGAAAACCATTTGAGGCACAGGCTCCTAAA TACGTACAACTAAATGTAGAAAATATTCAAGCTAATTCGAATGTTAAAGCAGGTATTTTTGCAGCATATAAAGCTATT GATTTCCATCCTCGATACAAGGATTATCTATTATTTGATAAAGAGAATATCAGTAAAGAAGATAGACAAAAGATTAAA GAACTTTCTTTGTCACAGGGATACGTTAAACTGCTAAATGCTTATCACAAAATCCCTGTTCTAGTCACGGGTTATGGC TATTCGACAGCGAGAGGTATTGCCCAAAAAGAAATTGATAAACGTCCTCTGCCGATTAATGAAAAAGAACAAGGTCAG CGTTTACTAGAAGATTATGAATCTTTTATATCATCCGGTAGTTTTGGAGCGACTATCAATGCATGGCAAGACGATTGG AATGCAAGGGCGTGGAATACATCCTTCGCCACAAATAAACATAGTCAATTCCTATGGGGGGGATGCACAAGTATTTAAT CAAGGTTATGGTTTATTAGGCTTTAAAAACGCAAAACATCATTATCAAGTTGATGGTAAAAGAGGCAAAGGAGAGTGG AAACATCCTCTG

### SEQ ID NO. 2802: SAG1552 FROM THE

### SEQ ID NO. 2803: SAG1552 FROM THE 18RS21 GBS TYPE II STRAIN

AAGGGCTTATTAAAAGAAAATACAAGAACTAACTTTGTTGTTAAAGGTGATACTGTACTTCACAAGCCCACCAATAAA CCTTTTGTTGTTAAAGGAGTAGACGTTGAGTCTTCCTTAGCGGGTTATCATCACAACGATTTTCCTATTACTCAAAAA ACGTATCGTGAATGGTTCCATTTAATTTCCAACATGGGGGCAAATACTGTAAGAGTCAAGGTACCGATGAATGTTGCA TTTTACGATGCCTTATATCACCACAACAAAGCATCAAAGAGGCCCACTGTATTTGTTGCAAGGAATACGTATAGATTCT TATCGCAATAATGCTTCTATAACAGCTTTTAATGATAATTATAGGGGGTATTTAAAACGAGAAGCAAAAGGCGTTGTG GATATTCTCCATGGGCGTAAGCAAGTATGGAATACTGATTTGGGTAGCCGTCATTATCATTATGATCTTAGTCCTTGG GTACTTGGTTATGTCGTAGGGGATGATTGGAATAGTGGTACTGTCGCTTATACTAATCATCAAGAGAAAAAAACGCAA TATAAAGGACGTTATTTTAAAACTTCTGTGGCAGCTAATCCATTTGAGGTCATGCTAGCTCAAGTAATGGATGAATTG ACACATTATGAGACAGCTAAATATGGTTGGCAACATTTGATTAGTTTTTCAAACTCACCAACAACAGACCCTTTTCAT TATCGAAAACCATTTGAGGCACAGGCTCCTAAATACGTACAACTAAATGTAGAAAATATTCAAGCTAATTCAAATGTT AAAGCAGGTATGTTTGCAGCATATAAAGCTATTGATTTCCATCCTCGATACAAGGATTATCTATTATTTGATAAAGAG CACAAAATCCCTGTTCTAGTCACGGGTTATGGCTATTCGACAGCGAGAGGTATTGCCCAAAAAGAAATTGATAAACGT CCTCTGCCGATTAATGAAAAAGAACAAGGTCAGCGTTTACTAGAAGATTATGAATCTTTATATATCATCCGGTAGTTTT GGAGCGACTATCAATGCATGGCAAGACGATTGGAATGCAAGGGCGTGGAATACATCTTTCGCCACAAATAAACATAGT CAATTCCTATGGGGGGATGCACAAGTATTTAATCAAGGTTATGGTTTATTAGGCTTTAAAAAACGCAAAACATCATTAT CAAGTTGATGGTAAAAGAGGCAAAGGAGAGTGGAAACATCCTCTGATGACTAGTGCAACAGGAGATGACTTATATGCT AGCAGTGATGAAAGCTATCTCTACCTTGCGATTAAAACAAAACCTGAAAAACTAAAAGAAAAACGATTATTACCAATA GATATTACACCAAAATCTGGTAGTAGAAAAATGAATGGTAGTAAGGTCACATTTTCTAAATCTAGTGACTTTGTATTG TCTATTGATCCAAATGGCAAGTCTGAATTATTTGTCCAAGAGCGCTATAATGCCTTAAAAGCGAACTATCTTCGACAG CTTAACGGTAAAGATTTTTATGCTTTCCCACCAAAGAAGAACAGTAGTAATTTTGAGCAGATAAATATGGTATTGAGA AATACAAAGATTGTTGAAGACATGGAAAAAGTAAAAGCAACAGAGAGGGTTCTTACCAACTCATCCTACTGGTCTTCTC AAAACAGGAACAACTGATAGGCACCAAAAAACATTTGATTCACAAACAGATATTTCGTTTGGAAAGGACTTTATAGAG GTCAGAATTCCGTGGCAGTTGTTGAATTTTTCTGATCCATCATCTCAAAAAATTCACGATGATTACTTTAAACATTAT GGTGTGAAGGAGTTAGAAATTGAGAGCATTGCTTTAGGATTAGGTGCTAATAGCAAAGAAAACACACTGATAAAGATG GCAGATTATCGTTTGAAAAATTGGGAGAGACCCGATACCAAAACCTTTTTAAAAGACTCCTATTATGTATTAAGAAAG AA



## (conserved hypothetical protein)

### SEQ ID NO. 2804: SAG1552 FROM THE 2603 V/R GBS TYPE V STRAIN (REVERSE COMPLEMENT)

TATTAAAAGAAATACAAGAACTAACTTTGTTGTTAAAGGTGATACTGTACTTCACAAGCCCACCAATAAACCTTTTG TTGTTAAAGGAGTAGACGTTGAGTCTTCCTTAGCGGGTTATCATCACAACGATTTTCCTATTACTCAAAAAACGTATC GTGAATGGTTCCATTTAATTTCCAACATGGGGGCAAATACTGTAAGAGTCAAGGTACCGATGAATGTTGCATTTTACG ATGCCTTATATCACCACAACAAAGCATCAAAGAGGCCACTGTATTTGTTGCAAGGAATACGTATAGATTCTTATCGCA ATAATGCTTCTATAACAGCTTTTAATGATAATTATAGGGGGTATTTAAAACGAGAAGCAAAAGGCGTTGTGGATATTC TCCATGGGCGTAAGCAAGTATGGAATACTGATTTGGGTAGCCGTCATTATCATTATGATCTTAGTCCTTGGGTACTTG GTTATGTCGTAGGGGATGATTGGAATAGTGGTACTGTCGCTTATACTAATCATCAAGAGAAAAAAACGCAATATAAAG GACGTTATTTTAAAACTTCTGTGGCAGCTAATCCATTTGAGGTCATGCTAGCTCAAGTAATGGATGAATTGACACATT ATGAGACAGCTAAATATGGTTGGCAACATTTGATTAGTTTTTCAAACTCACCAACAACAGACCCTTTTCATTATCGAA AACCATTTGAGGCACAGGCTCCTAAATACGTACAACTAAATGTAGAAAATATTCAAGCTAATTCAAATGTTAAAGCAG GTATGTTTGCAGCATATAAAGCTATTGATTTCCATCCTCGATACAAGGATTATCTATTATTTGATAAAGAGAATATCA GTAAAGAAGATAGACAAAAGATTAAAGAACTTTCTTTGTCACAGGGATACGTTAAACTGCTAAATGCTTATCACAAAA TÇCCTGTTCTAGTCACGGGTTATGGCTATTCGACAGCGAGAGGTATTGCCCCAAAAAGAAATTGATAAACGTCCTCTGC CGATTAATGAAAAAGAACAAGGTCAGCGTTTACTAGAAGATTATGAATCTTTTATATCATCCGGTAGTTTTGGAGCGA CTATCAATGCATGGCAAGACGATTGGAATGCAAGGGCGTGGAATACATCTTTCGCCACAAATAAACATAGTCAATTCC TATGGGGGGATGCACAAGTATTTAATCAAGGTTATGGTTTATTAGGCTTTAAAAACGCAAAACATCATTATCAAGTTG ATGGTAAAAGAGGCAAAGGAGAGTGGAAACATCCTCTGATGACTAGTGCAACAGGAGATGACTTATATGCTAGCAGTG ATGAAAGCTATCTCTACCTTGCGATTAAAACAAAACCTGAAAAACTAAAAGAAAACGATTATTACCAATAGATATTA CACCAAAATCTGGTAGTAGAAAATGAATGGTAGTAAGGTCACATTTTCTAAATCTAGTGACTTTGTATTGTCTATTG ATCCAAATGGCAAGTCTGAATTATTTGTCCAAGAGCGCTATAATGCCTTAAAAGCGAACTATCTTCGACAGCTTAACG GTAAAGATTTTTATGCTTTCCCACCAAAGAAGAACAGTAGTAATTTTGAGCAGATAAATATGGTATTGAGAAATACAA AGATTGTTGAAGACATGGAAAAAGTAAAAGCAACAGAGAGGTTCTTACCAACTCATCCTACTGGTCTTCTCAAAACAG GAACAACTGATAGGCACCAAAAAACATTTGATTCACAAACAGATATTTCGTTTGGAAAGGACTTTATAGAGGTCAGAA TTCCGTGGCAGTTGTTGAATTTTTCTGATCCATCATCTCAAAAAATTCACGATGATTACTTTAAACATTATGGTGTGA AGGAGTTAGAAATTGAGAGCATTGCTTTAGGATTAGGTGCTAATAGCAAAGAAACACACTGATAAAGATGGCAGATT CTAAAGAAAGAGAGAACATATGGTCCA

### SEQ ID NO. 2805: SAG1552 FROM THE A909 GBS TYPE Ia STRAIN (REVERSE COMPLEMENT)

AAGGGCTTATTAAAAGAAAATACAAGAACTAACTTTGTTGTTAAAGGTGATACTGTACTTCACAAGCCCACCAATAAA CCTTTTGTTGTTAAAGGAGTAGACGTTGAGTCTTCCTTAGCGGGTTATCATCACAACGATTTTCCTATTACTCAAAAA ACGTATCGTGAATGGTTCCATTTAATTTCCAACATGGGGGCAAATACTGTAAGAGTCAAGGTACCGATGAATGTTGCA TTTTACGATGCCTTATATCACCACAACAAAGCATCAAAGAGGCCACTGTATTTGTTGCAAGGAATACGTATAGATTCT TATCGCAATAATGCTTCTATAACAGCTTTTAATGATAATTATAGGGGGTATTTAAAACGAGAAGCCAAAAGGCGTTGTG GATATTCTCCATGGGCGTAAGCAAGTATGGAATACTGATTTGGGTAGCCGTCATTATCATTATGATCTTAGTCCTTGG GTACTTGGTTATGTCGTAGGGGATGATTGGAATAGTGGTACTGTCGCTTATACTAATCATCAAGAGAAAAAAACGCAA TATAAAGGACGTTATTTTAAAACTTCTGTGGCAGCTAATCCATTTGAGGTCATGCTAGCTCAAGTAATGGATGAATTG ACACATTATGAGACAGCTAAATATGGTTGGCAACATTTGATTAGTTTTTCAAACTCACCAACAACAGACCCTTTTCAT TATCGAAAACCATTTGAGGCACAGGCTCCTAAATACGTACAACTAAATGTAGAAAATATTCAAGCTAATTCAAATGTT **AAAGCAGGTATGTTTGCAGCATATAAAGCTATTGATTTCCATCCTCGATACAAGGATTATCTATTATTTGATAAAGAG** CACAAAATCCCTGTTCTAGTCACGGGTTATGGCTATTCGACAGCGAGAGGTATTGCCCAAAAAGAAATTGATAAACGT CCTCTGCCGATTAATGAAAAAGAACAAGGTCAGCGTTTACTAGAAGATTATGAATCTTTTATATCATCCGGTAGTTTT GGAGCGACTATCAATGCATGGCAAGACGATTGGAATGCAAGGGCGTGGAATACATCTTTCGCCACAAATAAACATAGT CAATTCCTATGGGGGGATGCACAAGTATTTAATCAAGGTTATGGTTTATTAGGCTTTAAAAAACGCAAAACATCATTAT CAAGTTGATGGTAAAAGAGGCAAAGGAGAGTGGAAACATCCTCTGATGACTAGTGCAACAGGAGATGACTTATATGCT AGCAGTGATGAAAGCTATCTCTACCTTGCGATTAAAACAAAACCTGAAAAACTAAAAGAAAAACGATTATTACCAATA GATATTACACCAAAATCTGGTAGTAGAAAAATGAATGGTAGTAAGGTCACATTTTCTAAATCTAGTGACTTTGTATTG TCTATTGATCCAAATGGCAAGTCTGAATTATTTGTCCAAGAGCGCTATAATGCCTTAAAAGCGAACTATCTTCGACAG CTTAACGGTAAAGATTTTTATGCTTTCCCACCAAAGAAGAACAGTAGTAATTTTGAGCAGATAAATATGGTATTGAGA AATACAAAGATTGTTGAAGACATGGAAAAAGTAAAAGCAACAGAGAGGTTCTTACCAACTCATCCTACTGGTCTTCTC AAAACAGGAACAACTGATAGGCACCAAAAAACATTTGATTCACAAACAGATATTTCGTTTGGAAAGGACTTTATAGAG GTCAGAATTCCGTGGCAGTTGTTGAATTTTTCTGATCCATCATCTCAAAGAATTCACGATGATTACTTTAAACATTAT GGTGTGAAGGAGTTAGAAAATTGAGAGCCATTGCTTTAGGATTAGGTGCTAATAGCAAAGAAAACACACTGATAAAGA TGGCAGATTATCGTTTGAAAAATTGGGAGAGACCCGATACCAAAACCTTTTTAAAAGA

# Table 28: C mparative Sequences relating to SAG1552 (conserved hypoth tical protein)

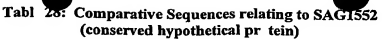
SEO ID NO. 2806: SAG1552 FROM THE CJB110 GBS NONTYPEABLE STRAIN GATACTGTACTTCACAAGCCCACCAATAAACCTTTTGTTGTTAAAGGAGTAGACGTTGAGTCTTCCTTAGCGGGTTAT CATCACAACGATTTTCCTATTACTCAAAAAACGTATCGTGAATGGTTCCATTTAATTTCCAACATGGGGGCAAATACT GTAAGAGTCAAGGTACCGATGAATGTTGCATTTTACGATGCCTTATATCACCACAACAAGCATCAAAGAGGCCACTG TATTTGTTGCAAGGAATACGTATAGATTCTTATCGCAATAATGCTTCTATAACAGCTTTTAATGATAATTATAGGGGG TATTTAAAACGAGAAGCAAAAGGCGTTGTGGATATTCTCCATGGGCGTAAGCAAGTATGGAATACAGATTTTGGTAGC CGTCATTATCATTATGATCTTAGTCCTTGGGTACTTGGTTATGTCGTAGGGGATGATTGGAATAGTGGTACTGTCGCT TATACTAATCATCAAGAGAAAAAAACGCAATATAAAGGACGTTATTTTAAAACTTCTGTGGCAGCTAATCCATTTGAG GTCATGCTAGCTCAAGTAATGGATGAATTGACACATTATGAGACAGCTAAATATGGTTGGCAACATTTGATTAGTTTT TCAAACTCACCAACAACAGACCCTTTTCATTATCGAAAACCATTTGAGGCACAGGCTCCTAAATACGTACAACTAAAT GTAGAAATATTCAAGCTAATTCAAATGTTAAAGCAGGTATGTTTGCAGCATATAAAGCTATTGATTTCCATCCTCGA CAGGGATACGTTAAACTGCTAAATGCTTATCACAAAATCCCTGTTCTAGTCACGGGTTATGGCTATTCGACAGCGAGA GGTATTGCCCAAAAAGAAATTGATAAACGTCCTCTGCCGATTAATGAAAAAGAACAAGGTCAGCGTTTACTAGAAGAT TATGAATCTTTATATCATCCGGTAGTTTTGGAGCGACTATCAATGCATGGCAAGACGATTGGAATGCAAGGGCGTGG AATACATCTTTCGCCACAAATAAACATAATCAATTCCTATGGGGGGATGCACAAGTATTTAATCAAGĞTTATGGTTTA TTAGGCTTTAAAAACGCAAAACATCATTATCAAGTTGATGGTAAAAGAGGCAAAGGAGAGTGGAAACATCCTCTGATG ACTAGTGCAACAGGAGATGACTTATATGCTAGCAGTGATGAAAGCTATCTCTACCTTGCGATTAAAACAAAACCTGAA ACATTTCTAAATCTAGTGACTTTGTATTGTCTATTGATCCAAATGGCAAGTCTGAATTATTTGTCCAAGAGCGCTAT AATGCCTTAAAAGCGAACTATCTTCGACAGCTTAACGGTAAAGATTTTTATGCTTTCCCACCAAAGAAGAACAGTAGT **AATTTTGAGCAGATAAATATGGTATTGAGAAATACAAAGATTGTTGAAGACATGGAAAAAGTAAAAGCAACAGAGAG** TTCTTACCAACTCATCCTACTGGTCTTCTCAAAACAGGAACAACTGATAGGCACCAAAAAACATTTGATTCACAAACA GATATTTCGTTTGGAAAGGACTTTATAGAGGTCAGAATTCCGTGGCAGTTGTTGAATTTTTCTGATCCATCATCTCAA **AAAATTCACGATGATTACTTTAAACATTATGGTGTGAAGGAGTTAGAAATTGAGAGCATTGCTTTAGGATTAGGTGCT AATAGCAAAGAAAACACTGATAAAGATGGCAGATTATCGTTTGAAAAATTGGGAGAGACCCGATACCAAAACCTTT** TTAAAAGACTCCTATTATGTATTAAGAAAGA

SEQ ID NO. 2807: SAG1552 FROM THE COH1 GBS TYPE III STRAIN CAATAAACCTTTTGTTGTTAAAGGAGTAGACGTTGAGTCTTCCTTAGCGGGTTATCATCACAACGATTTTCCTATTAC TCAAAAAACGTATCGTGAATGGTTCCATTTAATTTCCAACATGGGGGCAAATACTGTAAGAGTCAAGGTACCGATGAA TGTTGCATTTTACGATGCCTTATATCACCACAACAAGAATCAAAGAGCCCACTGTATTTGTTGCAAGGAATACGTAT AGATTCTTATCGCAATAATGCTTCTATAACAGCTTTTAATGATAATTATAGGGGGTTATTAAAACGAGAAGCAAAAGG CGTTGTGGATATTCTCCATGGGCGTAAGCAAGTATGGAATACTGATTTTGGTAGCCGTCATTATCATTATGATCTTAG TCCTTGGGTACTTGGTTATGTCGTAGGGGATGATTGGAATAGTGGTACTGTCGCTTATACTAATCATCAAGAGAAAAA AACGCAATATAAAGGACGTTATTTTAAAACTTCTGTGGCAGCTAATCCATTTGAGGTCATGCTAGCTCAAGTAATGGA TGAATTGACACATTATGAGACAGCTAAATATGGTTGGCAACATTTGATTAGTTTTTCAAACTCACCAACAACAGACCC TTTTCATTATCGAAAACCATTTGAGGCACAGGCTCCTAAATACGTACAACTAAATGTAGAAAATATTCAAGCTAATTC AAATGTTAAAGCAGGTATGTTTGCAGCATATAAAGCTATTGATTTCCATCCTCGATACAAGGATTATCTATTATTTGA TGCTTATCACAAAATCCCTGTTCTAGTCACGGGTTATGGCTATTCGACAGCGAGAGGTATTGCCCCAAAAAGAAATTGA TAAACGTCCTCTGCCGATTAATGAAAAAGAACAAGGTCAGCGTTTACTAGAAGATTATGAATCTTTTATATCATCCGG TAGTTTTGGAGCGACTATCAATGCATGGCAAGACGATTGGAATGCAAGGGCGTGGAATACATCTTTCGCCACAAATAA **ACATAGTCAATTCCTATGGGGGGATGCACAAGTATTTAATCAAGGTTATGGTTTATTAGGCTTTAAAAAACGCAAAACA** TCATTATCAAGTTGATGGTAAAAGAGGCAAAGGAGAGTGGAAACATCCTCTGATGACTAGTGCAACAGGAGATGACTT ATATGCTAGCAGTGATGAAAGCTATCTCTACCTTGCGATTAAAACAAAACCTGAAAAACTAAAAGAAAACGATTATT TGTATTGTCTATTGATCCAAATGGCAAGTCTGAATTATTTGTCCAAGAGCGCTATAATGCCTTAAAAGCGAACTATCT TCGACAGCTTAACGGTAAAGATTTTTATGCTTTCCCACCAAAGAAGAACAGTAGTAATTTTGAGCAGATAAATATGGT ATTGAGAAATACAAAGATTGTTGAAGACATGGAAAAAGTAAAAGCAACAGAGAGGTTCTTACCAACTCATCCTACTGG TCTTCTCAAAACAGGAACAACTGATAGGCACCAAAAAACATTTGATTCACAACCAGATATTTCGTTTGGAAAGGACTT TATAGAGGTCAGAATTCCGTGGCAGTTGTTGAATTTTTCTGATCCATCATCTCAAAAAATTCACGATGATTACTTTAA ACATTATGGTGTGAAGGAGTTAGAAATTGAGAGCATTGCTTTAGGATTAGGTGCTAATAGCAAAGAAAACACACTGAT AAAGATGGCAGATTATCGTTTGAAAAATTGGGAGAGACCCGATACCAAAACCTTTTTAAAAGACT

## Table 28: Comparative Sequences relating to SAG1552 (conserved hypothetical pr tein)

SAG1552 FROM THE H36b GBS TYPE Ib STRAIN SEQ ID NO. 2808: AAGGGGCTTATTAAAAGAAAATACAAGAACTAACTTTGTTGTTAAAGGTGATACTGTACTTCACAAGCCCACCAATAA ACCTTTTGTTGTTAAAGGAGTAGACGTTGAGTCTTCCTTAGCGGGTTATCATCACAACGATTTTCCTATTACTCAAAA AACGTATCGTGAATGGTTCCATTTAATTTCCAACATGGGGGCAAATACTGTAAGAGTCAAGGTACCGATGAATGTTGC ATTTTACGATGCCTTATATCACCACAACAAGCATCAAAGAGGCCACTGTATTTGTTGCAAGGAATACGTATAGATTC TTATCGCAATAATGCTTCTATAACAGCTTTTAATGATAATTATAGGGGGTATTTAAAACGAGAAGCAAAAGGCGTTGT GGATATTCTCCATGGGCGTAAGCAAGTATGGAATACTGATTTTGGTAGCAGTCATTATCATTATGATCTTAGTCCTTG GGTACTTGGTTATGTCGTAGGGGATGATGGACATAGTGGTACTGTCGCTTTATACTAATCATCAAGAGGAGAAAAAACG CAATATAAAGGACGTTATTTTAAAACTTCTGTGGCAGCTAATCCATTTGAGGTCATGCTAGCTCAAGTAATGGATGAA TTGACACATTATGAGACAGCTAAATATGGTTGGCAACATTTGATTAGTTTTTCAAACTCACCAACAACAGACCCTTTT CATTATCGAAAACCATTTGAGGCACAGGCTCCTAAATACGTACAACTAAATGTAGAAAATATTCAAGCTAATTCGAAT GTTAAAGCAGGTATGTTTGCAGCATATAAAGCTATTGATTTCCATCCTCGATACAAGGATTATCTATTATTTGATAAA TATCACAAAATCCCTGTTCTAGTCACGGGTTATGGCTACTCGACAGCGAGAGGTATTGCCCAAAAAGAAATTGATAAA CGTCCTCTGCCGATTAATGAAAAAGAACAAGGTCAGCGTTTACTAGAAGATTATGAATCTTTTATATCATCCGGTAGT TTTGGAGCGACTATCAATGCATGGCAAGACGATTGGAATGCAAGGGTGTGGAATACATCCTTCGCCACAAATAAACAT AGTCAATTCCTATGGGGGGATGCACAAGTATTTAATCAAGGTTATGGTTTATTAGGCTTTAAAAAACGCAAAACATCAT TATCAGGTTGATGGTAAAAGAGGCAAAGAAGAGTGGAAACATCCTCTGATGACTAGTGCAACAGGAGATGACTTATAT GCTAGCAGTGATGAAAGCTATCTCTACCTTGCGATTAAAACAAAACCTGAAAAACTAAAAGAAAAACGATTATTACCA TTGTCTATTGATCCAAATGGCAAGTCTGAATTATTTGTCCAAGAGCGCTATAACGCCTTAAAAGCGAACTATCTTCGA CAGCTTAATGGTAAAGATTTTTATGCTTTCCCACCAAAGAAGAACAGTAGTAATTTTGAGCAGATAAATATGGTATTG AGAAATACAAAGATTGTTGAAGACATGGAAAAAGTAAAAGCAACAGAGAGGTTCTTACCAACTCATCCTACTGGTCTT CTCAAAACAGGAACAACTGATAGGCACCAAAAAACATTTGATTCACAAACAGATATTTCGTTTGGAAAGGACTTTATA GAGGTCAGAATTCCGTGGCAGTTGTTGAATTTTTCTGATCCATCATCTCAAAAAATTCACGATGATTACTTTAAACAT TATGGTGTGAAGGAGTTAGAAATTGAGAGCATTGCTTTAGGATTAGGTGCTAATAGCAAAGAAAACACACTGATAAAG ATGGCAGATTATCGTTTGAAAAATTGGGAGAGACCCGATACCAAAACCTTTTTAAAAAGACTCCTATTATAGT

SEQ ID NO. 2809: SAG1552 FROM THE JM9130013 GBS TYPE VIII STRAIN **ACTTTGTTGTTAAAGGTGATACTGTACTTCACAAGCCCACCAATAAACCTTTTGTTGTTAAAGGAGTAGACGTTGAGT** CTTCCTTAGCGGGTTATCATCACAACGATTTTCCTATTACTCAAAAAACGTATCGTGAATGGTTCCATTTAATTTCCA ACATGGGGGCAAATACTGTAAGAGTCAAGGTACCGATGAATGTTGCATTTTACGATGCCTTATATCACCACAACAAAG CATCAAAGAGGCCACTGTATTTGTTGCAAGGAATACGTATAGATTCTTATCGCAATAATGCTTCTATAACAGCTTTTA ATGATAATTATAGGGGGTATTTAAAACGAGAAGCAAAAGGCGTTGTGGATATTCTCCATGGGCGTAAGCAAGTATGGA ATACTGATTTTGGTAGCAGTCATTATCATTATGATCTTAGTCCTTGGGTACTTGGTTATGTCGTAGGGGATGATTGGA ATAGTGGTACTGTCGCTTATACTAATCATCAAGAGAAAAAAACGCAATATAAAGGACGTTATTTTAAAACTTCTGTGG CAGCTAATCCATTTGAGGTCATGCTAGCTCAAGTAATGGATGAATTGACACATTATGAGACAGCTAAATATGGTTGGC AACATTTGATTAGTTTTTCAAACTCACCAACAACAGACCCTTTTCATTATCGAAAACCATTTGAGGCACAGGCTCCTA AATACGTACAACTAAATGTAGAAAATATTCAAGCTAATTCGAATGTTAAAGCAGGTATGTTTGCAGCATATAAAGCTA TTGATTTCCATCCTCGATACAAGGATTATCTATTATTTGATAAAGAGAATATCAGTAAAGAAGATAGACAAAAGATTA AAGAACTTTCTTTGTCACAGGGATACGTTAAACTGCTAAATGCTTATCACAAAATCCCTGTTCTAGTCACGGGTTATG GCTACTCGACAGCGAGAGGTATTGCCCAAAAAGAAATTGATAAACGTCCTCTGCCGATTAATGAAAAAGAACAAGGTC AGCGTTTACTAGAAGATTATGAATCTTTTATATCATCCGGTAGTTTTGGAGCGACTATCAATGCATGGCAAGACGATT GGAATGCAAGGGTGTGGAATACATCCTTCGCCACAAATAAACATAGTCAATTCCTATGGGGGGGATGCACAAGTATTTA ATCAAGGTTATGGTTTATTAGGCTTTAAAAACGCAAAACATCATTATCAGGTTGATGGTAAAAAGAGGCAAAGAAGAGT GGAAACATCCTCTGATGACTAGTGCAACAGGAGATGACTTATATGCTAGCAGTGATGAAAGCTATCTCTACCTTGCGA TTAAAACAAAACCTGAAAAACTAAAAGAAAAACGATTATTACCAATAGATATTACACCAAAATCTGGTAGTAGAAAAA TGAATGGTAGTAAGGTCACATTTTCTAAATCTAGTGACTTTGTATTGTCTATTGATCCAAATGGCAAGTCTGAATTAT  ${\tt TTGTCCAAGAGCGCTATAACGCCTTAAAAGCGAACTATCTTCGACAGCTTAATGGTAAAGATTTTTATGCTTTCCCAC}$ CAAAGAAGAACAGTAGTAATTTTGAGCAGATAAATATGGTATTGAGAAATACAAAGATTGTTGAAGACATGGAAAAAAG TAAAAGCAACAGAGAGGTTCTTACCAACTCATCCTACTGGTCTTCTCAAAACAGGAACAACTGATAGGCACCAAAAAA CATTTGATTCACAAACAGATATTTCGTTTGGAAAGGACTTTATAGAGGTCAGAATTCCGTGGCAGTTGTTGAATTTTT CTGATCCATCATCACAAAAAATTCACGATGATTACTTTAAACATTATGGTGTGAAGGAGTTAGAAATTGAGAGCATTG CTTTAGGATTAGGTGCTAATAGCAAAGAAAACACACTGATAAAGATGGCAGATTATCGTTTGAAAAAATTGGGAGAGAC CCGATACCAAAACCTTTTTAAAAGACTCCTATTATAGTATTAAGAAAG



SEQ ID NO. 2810: SAG1552 FROM THE M732 GBS TYPE III STRAIN TACAAGAACTAACTTTGTTGTTAAAGGTGATACTGTACTTCACAAGCCCACCAATAAACCTTTTGTTGTTAAAGGAGT AGACGTTGAGTCTTCCTTAGCGGGTTATCATCACAACGATTTTCCTATTACTCAAAAAACGTATCGTGAATGGTTCCA TTTAATTTCCAACATGGGGGCAAATACTGTAAGAGTCAAGGTACCGATGAATGTTGCATTTTACGATGCCTTATATCA CCACAACAAAGAATCAAAGAGGCCACTGTATTTGTTGCAAGGAATACGTATAGATTCTTATCGCAATAATGCTTCTAT AACAGCTTTTAATGATAATTATAGGGGGTATTTAAAACGAGAAGCAAAAGGCGTTGTGGATATTCTCCATGGGCGTAA GCAAGTATGGAATACTGATTTTGGTAGCCGTCATTATCATTATGATCTTAGTCCTTGGGTACTTGGTTATGTCGTAGG GGATGATTGCAATAGTGGTACTGTCGCTTATACTAATCATCAAGAGAAAAAAACGCAATATAAAGGACGTTATTTTAA AACTTCTGTGGCAGCTAATCCATTTGAGGTCATGCTAGCTCAAGTAATGGATGAATTGACACATTATGAGACAGCTAA ATATGGTTGGCAACATTTGATTATTTCAAACTCACCAACAACAGACCCTTTTCATTATCGAAAACCATTTGAGGC ACAGGCTCCTAAATACGTACAACTAAATGTAGAAAATATTCAAGCTAATTCAAATGTTAAAGCAGGTATGTTTGCAGC ATATAAAGCTATTGATTTCCATCCTCGATACAAGGATTATCTATTATTTGATAAAGAGAATATCAGTAAAGAAGATAG ACAAAAGATTAAAGAACTTTCTTTGTCACAGGGATACGTTAAACTGCTAAATGCTTATCACAAAATCCCTGTTCTAGT CACGGGTTATGGCTATTCGACAGCGAGAGGTATTGCCCCAAAAAGAAATTGATAAACGTCCTCTGCCGATTAATGAAAA AGAACAAGGTCAGCGTTTACTAGAAGATTATGAATCTTTTATATCATCCGGTAGTTTTGGAGCGACTATCAATGCATG GCAAGACGATTGGAATGCAAGGGCGTGGAATACATCTTTCGCCACAAATAAACATAGTCAATTCCTATGGGGGGATGC ACAAGTATTTAATCAAGGTTATGGTTTATTAGGCTTTAAAAACGCAAAACATCATTATCAAGTTGATGGTAAAAGAGG CAAAGGAGAGTGGAAACATCCTCTGATGACTAGTGCAACAGGAGATGACTTATATGCTAGCAGTGATGAAAGCTATCT CTACCTTGCGATTAAAACCAAAACCTGAAAAACTAAAAGAAAAACGATTATTACCAATAGATATTACACCAAAATCTGG TAGTAGAAAAATGAATGGTAGTAAGGTCACATTTTCTAAATCTAGTGACTTTGTATTGTCTATTGATCCAAATGGCAA GTCTGAATTATTTGTCCAAGAGCGCTATAATGCCTTAAAAGCGAACTATCTTCGACAGCTTAACGGTAAAGATTTTTA TGCTTTCCCACCAAAGAAGAACAGTAGTAATTTTGAGCAGATAAATATGGTATTGAGAAATACAAAGATTGTTGAAGA CATGGAAAAAGTAAAAGCAACAGAGAGGTTCTTACCAACTCATCCTACTGGTCTTCTCAAAACAGGAACAACTGATAG GCACCAAAAAACATTTGATTCACAAACAGATATTTCGTTTGGAAAGGACTTTATAGAGGTCAGAATTCCGTGGCAGTT GTTGAATTTTTCTGATCCATCATCTCAAAAAATTCACGATGATTACTTTAAACATTATGGTGTGAAGGAGTTAGAAAT TGAGAGCATTGCTTTAGGATTAGGTGCTAATAGCAAAGAAACACACTGATAAAGATGGCAGATTATCGTTTGAAAAA TTGGGAGAGACCCGATACCAAAACCTTTTTAAAAGACTCCTATTATAGTATTAAG

### SEQ ID NO. 2811: SAG1552 FROM THE M781 GBS TYPE III STRAIN GTACTTCACAAGCCCACCAATAAACCTTTTGTTGTTAAAGGAGTAGACGTTGAGTCTTCCTTAGCGGGTTATCATCAC AACGATTTTCCTATTACTCAAAAAACGTATCGTGAATGGTTCCATTTAATTTCCAACATGGGGGCAAATACTGTAAGA GTCAAGGTACCGATGAATGTTGCATTTTACGATGCCTTATATCACCACAACAAAGAATCAAAGAGGCCACTGTATTTG TTGCAAGGAATACGTATAGATTCTTATCGCAATAATGCTTCTATAACAGCTTTTAATGATAATTATAGGGGGTATTTA AAACGAGAAGCAAAAGGCGTTGTGGATATTCTCCATGGGCGTAAGCAAGTATGGAATACTGATTTTGGTAGCCGTCAT TATCATTATGATCTTAGTCCTTGGGTACTTGGTTATGTCGTAGGGGATGATTGGAATAGTGGTACTGTCGCTTATACT AATCATCAAGAGAAAAAAACGCAATATAAAGGACGTTATTTTAAAACTTCTGTGGCAGCTAATCCATTTGAGGTCATG CTAGCTCAAGTAATGGATGAATTGACACATTATGAGACAGCTAAATATGGTTGGCAACATTTGATTAGTTTTTCAAAC TCACCAACAACAGACCCTTTTCATTATCGAAAACCATTTGAGGCACAGGCTCCTAAATACGTACAACTAAATGTAGAA AATATTCAAGCTAATTCAAATGTTAAAGCAGGTATGTTTGCAGCATATAAAGCTATTGATTTCCATCCTCGATACAAG TACGTTAAACTGCTAAATGCTTATCACAAAATCCCTGTTCTAGTCACGGGTTATGGCTATTCGACAGCGAGAGGTATT GCCCAAAAAGAAATTGATAAACGTCCTCTGCCGATTAATGAAAAAGAACAAGGTCAGCGTTTACTAGAAGATTATGAA TCTTTTATATCATCCGGTAGTTTTGGAGCGACTATCAATGCATGGCAAGACGATTGGAATGCAAGGGCGTGGAATACA TCTTTCGCCACAAATAAACATAGTCAATTCCTATGGGGGGATGCACAAGTATTTAATCAAGGTTATGGTTTATTAGGC TTTAAAAACGCAAAACATCATTATCAAGTTGATGGTAAAAGAGGCAAAGGAGAGTGGAAACATCCTCTGATGACTAGT GCAACAGGAGATGACTTATATGCTAGCAGTGATGAAAGCTATCTCTACCTTGCGATTAAAACAAAACCTGAAAAACTA TCTAAATCTAGTGACTTTGTATTGTCTATTGATCCAAATGGCAAGTCTGAATTATTTGTCCAAGAGCGCTATAATGCC TTAAAAGCGAACTATCTTCGACAGCTTAACGGTAAAGATTTTTATGCTTTCCCACCAAAGAAGAACAGTAGTAATTTT GAGCAGATAAATATGGTATTGAGAAATACAAAGATTGTTGAAGACATGGAAAAAGTAAAAGCAACAGAGAGGTTCTTA CCAACTCATCCTACTGGTCTTCTCAAAACAGGAACAACTGATAGGCACCAAAAAACATTTGATTCACAAACAGATATT TCGTTTGGAAAGGACTTTATAGAGGTCAGAATTCCGTGGCAGTTGTTGAATTTTTCTGATCCATCATCTCAAAAAATT CACGATGATTACTTTAAACATTATGGTGTGAAGGAGTTAGAAATTGAGAGCATTGCTTTAGGATTAGGTGCTAATAGC AAAGAAAACACACTGATAAAGATGGCAGATTATCGTTTGAAAAATTGGGAGAGACCCGATACCAAAACCTTTTTAAAA

GACTCCTATTATAGTATTAAGAAAGAATGG



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# Table 28: Comparative Sequences relating to SAG1552 (conserved hyp thetical protein)

6700001	
SEQ2801 SEQ2802	
SEQ2803	AAGGGCTTATTAAAAGAAAATACAAGAACT
SEQ2804	TATTAAAAGAAATACAAGAACT
SEQ2805	AAGGGCTTATTAAAAGAAATACAAGAACT
SEQ2806	ATTACTTTGATGGTAGTTTGTATTTACCAAAGGGCTTATTAAAAGAAAATACAAGAACT
SEQ2807	TTTACCACAGGGCTTATTAAAAGAAAATACAAGAACT
SEQ2808	AAGGGGCTTATTAAAAGAAAATACAAGAACT
SEQ2809	
SEQ2810	TACAAGAACT
SEQ2811	TTTGATGGTAGTTTGTATTTACCACAGGGCTTATTAAAAGAAAATACAAGAACT
SEQ2801 SEQ2802	TTTGTTGTTAAAGGTGATACTGTACTTCACAAGCCCACCAATAAACCTTTTGTTGTT
SEQ2803	ACTTTGTTGTTAAAGGTGATACTGTACTTCACAAGCCCACCAATAAACCTTTTGTTGTT
SEQ2804	ACTTTGTTGTTAAAGGTGATACTGTACTTCACAAGCCCACCAATAAACCTTTTGTTGTT
SEQ2805	ACTTTGTTGTTAAAGGTGATACTGTACTTCACAAGCCCACCAATAAACCTTTTGTTGTT
SEQ2806	ACTTTGTTGTTAAAGGTGATACTGTACTTCACAAGCCCACCAATAAACCTTTTGTTGTT
SEQ2807	ACTTTGTTGTTAAAGGTGATACTGTACTTCACAAGCCCACCAATAAACCTTTTGTTGTT
SEQ2808	ACTTTGTTGTTAAAGGTGATACTGTACTTCACAAGCCCACCAATAAACCTTTTGTTGTT
SEQ2809	ACTTTGTTGTTAAAGGTGATACTGTACTTCACAAGCCCACCAATAAACCTTTTGTTGTT
SEQ2810	ACTTTGTTGAAAGGTGATACTGTACTTCACAAGCCCACCAATAAACCTTTTGTTGTT
SEQ2811	ACTTTGTTGTTAAAGGTGATACTGTACTTCACAAGCCCACCAATAAACCTTTTGTTGTT
SEQ2801	AAGGAGTAGACGTTGAGTCTTCCTTAGCAGGTTATCATCACAACGATTTTCCTATTACT
SEQ2802	AAGGAGIAGACGIIGAGICIICCIIAGCAGGIIAICACAACGAIIITCCTATTACT
SEQ2803	AAGGAGTAGACGTTGAGTCTTCCTTAGCGGGTTATCATCACAACGATTTTCCTATTACT
SEQ2804	AAGGAGTAGACGTTGAGTCTTCCTTAGCGGGTTATCACACAACGATTTTCCTATTACT
SEQ2805	AAGGAGTAGACGTTGAGTCTTCCTTAGCGGGTTATCATCACAACGATTTTCCTATTACT
SEQ2806	AAGGAGTAGACGTTGAGTCTTCCTTAGCGGGTTATCATCACAACGATTTTCCTATTACT
SEQ2807	AAGGAGTAGACGTTGAGTCTTCCTTAGCGGGTTATCATCACAACGATTTTCCTATTACT
SEQ2808	AAGGAGTAGACGTTGAGTCTTCCTTAGCGGGTTATCATCACAACGATTTTCCTATTACT
SEQ2809	AAGGAGTAGACGTTGAGTCTTCCTTAGCGGGTTATCATCACAACGATTTTCCTATTACT
SEQ2810	AAGGAGTAGACGTTGAGTCTTCCTTAGCGGGTTATCATCACAACGATTTTCCTATTACT
SEQ2811	AAGGAGTAGACGTTGAGTCTTCCTTAGCGGGTTATCATCACAACGATTTTCCTATTACT
SEQ2801	AAAAAACGTATCGTGAGTGGTTCCATTTAATTTCCAACATGGGGGCAAATACTGTAAGA
SEQ2802	
SEQ2803	AAAAAACGTATCGTGAATGGTTCCATTTAATTTCCAACATGGGGGCAAATACTGTAAGA
SEQ2804	AAAAAACGTATCGTGAATGGTTCCATTTAATTTCCAACATGGGGGCAAATACTGTAAGA
SEQ2805	AAAAACGTATCGTGAATGGTTCCATTTAATTTCCAACATGGGGGCAAATACTGTAAGA
SEQ2806	AAAAAACGTATCGTGAATGGTTCCATTTAATTTCCAACATGGGGGCAAATACTGTAAGA
SEQ2807	AAAAAACGTATCGTGAATGGTTCCATTTAATTTCCAACATGGGGGCAAATACTGTAAGA
SEQ2808	AAAAAACGTATCGTGAATGGTTCCATTTAATTTCCAACATGGGGGCAAATACTGTAAGA
SEQ2809	AAAAAACGTATCGTGAATGGTTCCATTTAATTTCCAACATGGGGGCAAATACTGTAAGA
SEQ2810	AAAAAACGTATCGTGAATGGTTCCATTTAATTTCCAACATGGGGGCAAATACTGTAAGA
SEQ2811	AAAAAACGTATCGTGAATGGTTCCATTTAATTTCCAACATGGGGGCAAATACTGTAAGA
SEQ2801	TCAAAGTACCGATGAATGTTGCATTTTACGATGCTTTATATCACCACAACAAAGCATCA
SEQ2802	
SEQ2803	TCAAGGTACCGATGAATGTTGCATTTTACGATGCCTTATATCACCACAACAAAGCATCA
SEQ2804	TCAAGGTACCGATGAATGTTGCATTTTACGATGCCTTATATCACCACAACAAGCATCA
SEQ2805	TCAAGGTACCGATGAATGTTGCATTTTACGATGCCTTATATCACCACAACAAGCATCA
SEQ2806 SEQ2807	TCAAGGTACCGATGAATGTTGCATTTTACGATGCCTTATATCACCACAACAAGCATCA
SEQ2808	TCAAGGTACCGATGAATGTTGCATTTTACGATGCCTTATATCACCACAACAAGAATCA
SEQ2809	TCAAGGTACCGATGAATGTTGCATTTTACGATGCCTTATATCACCACAACAAGCATCA TCAAGGTACCGATGAATGTTGCATTTTACGATGCCTTATATCACCACAACAAGCATCA
SEQ2810	TCAAGGTACCGATGAATGTTGCATTTTACGATGCCTTATATCACCACAACAAGCATCA TCAAGGTACCGATGAATGTTGCATTTTACGATGCCTTATATCACCACAACAAGCATCA
SEQ2811	TCAAGGTACCGATGAATGTTGCATTTTACGATGCCTTATATCACCACAACAAGAATCA TCAAGGTACCGATGAATGTTGCATTTTACGATGCCTTATATCACCACAACAAGAATCA
<b></b>	
SEQ2801	AGAGGCCACTGTATTTGTTGCAAGGAATACGTATAGATTCTTATCGCAATAATGCTTCT
SEQ2802	
SEQ2803	AGAGGCCACTGTATTTGTTGCAAGGAATACGTATAGATTCTTATCGCAATAATGCTTCT
SEQ2804	AGAGGCCACTGTATTTGTTGCAAGGAATACGTATAGATTCTTATCGCAATAATGCTTCT
SEQ2805	AGAGGCCACTGTATTTGTTGCAAGGAATACGTATAGATTCTTATCGCAATAATGCTTCT
SEQ2806	AGAGGCCACTGTATTTGTTGCAAGGAATACGTATAGATTCTTATCGCAATAATGCTTCT
SEQ2807	AGAGGCCACTGTATTTGTTGCAAGGAATACGTATAGATTCTTATCGCAATAATGCTTCT
SEQ2808	AGAGGCCACTGTATTTGTTGCAAGGAATACGTATAGATTCTTATCGCAATAATGCTTCT



# Table 28: Comparative Sequences relating to SAG1552 (conserved hypothetical protein)

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SEQ2809	AGAGGCCACTGTATTTGTTGCAAGGAATACGTATAGATTCTTATCGCAATAATGCTTCT
SEQ2810	AGAGGCCACTGTATTTGTTGCAAGGAATACGTATAGATTCTTATCGCAATAATGCTTCT
SEQ2811	AGAGGCCACTGTATTTGTTGCAAGGAATACGTATAGATTCTTATCGCAATAATGCTTCT
SEQ2801	
	TAACAGCTTTTAATGATAATTATAGGGGGTATTTAAAACGAGAAGCAAAAGGCGTTGTG
SEQ2802	
SEQ2803	TAACAGCTTTTAATGATAATTATAGGGGGTATTTAAAACGAGAAGCAAAAGGCGTTGTG
_	THOOGET ITTATEMITATION THE TATABET OF THE TATABET O
SEQ2804	TAACAGCTTTTAATGATAATTATAGGGGGTATTTAAAACGAGAAGCAAAAGGCGTTGTG
SEQ2805	TAACAGCTTTTAATGATAATTATAGGGGGTATTTAAAACGAGAAGCAAAAGGCGTTGTG
<del></del>	THE STORY OF THE S
SEQ2806	TAACAGCTTTTAATGATAATTATAGGGGGTATTTAAAACGAGAAGCAAAAGGCGTTGTG
SEQ2807	TAACAGCTTTTAATGATAATTATAGGGGGTATTTAAAACGAGAAGCAAAAGGCGTTGTG
SEQ2808	
_	TAACAGCTTTTAATGATAATTATAGGGGGTATTTAAAACGAGAAGCAAAAGGCGTTGTG
SEQ2809	TAACAGCTTTTAATGATAATTATAGGGGGTATTTAAAACGAGAAGCAAAAGGCGTTGTG
SEQ2810	TAACAGCTTTTAATGATAATTATAGGGGGTATTTAAAACGAGAAGCAAAAGGCGTTGTG
	TAACAGCTTTTAATGATAATTATAGGGGGTATTTAAAACGAGAAGCAAAAGGCGTTGTG
SEQ2811	TAACAGCTTTTAATGATAATTATAGGGGGTATTTAAAACGAGAAGCAAAAGGCGTTGTG
0700001	
SEQ2801	ATATTCTCCATGGGCGTAAGCAAGTATGGAATACTGATTTTGGTAGCCGTCATTATCAT
SEQ2802	ATGACTA-GTGCAACAGGAGATGACTTATAT-GCTAGCAGTGATGAAAGC
	A WAR WOOD CONTROL OF THE CONTROL OF
SEQ2803	ATATTCTCCATGGGCGTAAGCAAGTATGGAATACTGATTTGGGTAGCCGTCATTATCAT
SEQ2804	ATATTCTCCATGGGCGTAAGCAAGTATGGAATACTGATTTGGGTAGCCGTCATTATCAT
SEQ2805	A TATA TO CONTROL CONTROL AND CONTROL AND CONTROL CONT
	ATATTCTCCATGGGCGTAAGCAAGTATGGAATACTGATTTGGGTAGCCGTCATTATCAT
SEQ2806	ATATTCTCCATGGGCGTAAGCAAGTATGGAATACAGATTTTGGTAGCCGTCATTATCAT
SEQ2807	ATATTCTCCATGGGCGTAAGCAAGTATGGAATACTGATTTTGGTAGCCGTCATTATCAT
	THE PROPERTY OF THE PROPERTY O
SEQ2808	ATATTCTCCATGGGCGTAAGCAAGTATGGAATACTGATTTTGGTAGCAGTCATTATCAT
SEQ2809	ATATTCTCCATGGGCGTAAGCAAGTATGGAATACTGATTTTGGTAGCAGTCATTATCAT
SEQ2810	ATA THOUGO A TOO COOK A COAR A
	ATATTCTCCATGGGCGTAAGCAAGTATGGAATACTGATTTTGGTAGCCGTCATTATCAT
SEQ2811	ATATTCTCCATGGGCGTAAGCAAGTATGGAATACTGATTTTGGTAGCCGTCATTATCAT
,	
SEQ2801	TATGATCTTAGTCCTTGGGTACTTGGTTATGTCGTAGGGGATGATTGGAATAGTGGT-AC
SEQ2802	TATCTCTACCTTGCG-ATTAAAACAAAACCTGAAAAACTAAAAGAAAAACGATTAT
SEQ2803	MACON COMPACTOR CONTROL OF CONTRO
	TATGATCTTAGTCCTTGGGTACTTGGTTATGTCGTAGGGGATGATTGGAATAGTGGT-AC
SEQ2804	TATGATCTTAGTCCTTGGGTACTTGGTTATGTCGTAGGGGATGATTGGAATAGTGGT-AC
SEQ2805	TATGATCTTAGTCCTTGGGTACTTGGTTATGTCGTAGGGGATGATTGGAATAGTGGT-AC
_	TATOMICTIAGICCI IGGILACI IGGILAGICGIAGGGATGAT TGGAATAGTGGT-AC
SEQ2806	TATGATCTTAGTCCTTGGGTACTTGGTTATGTCGTAGGGGATGATTGGAATAGTGGT-AC
SEQ2807	TATGATCTTAGTCCTTGGGTACTTGGTTATGTCGTAGGGGATGATTGGAATAGTGGT-AC
· SEQ2808	MARCA MORNA COROLLA CO
_	TATGATCTTAGTCCTTGGGTACTTGGTTATGTCGTAGGGGATGATGGACATAGTGGT-AC
·SEQ2809	TATGATCTTAGTCCTTGGGTACTTGGTTATGTCGTAGGGGATGATTGGAATAGTGGT-AC
SEQ2810	TATGATCTTAGTCCTTGGGTACTTGGTTATGTCGTAGGGGATGATTGCAATAGTGGT-AC
	TATAL COLLEGE ACTION INTO COLLEGE ACTION OF THE COLLEGE ACTION OF
SEQ2811	TATGATCTTAGTCCTTGGGTACTTGGTTATGTCGTAGGGGATGATTGGAATAGTGGT-AC
SEQ2801	PCPCCCPP ADACMANCARCARCA CARA ARRANGA CARA CARA CARA CARA CARA CARA CARA CA
	TGTCGCTT-ATACTAATCATCAAGAGAA-AAAAACGCAATATAAAGGAC-GTTATTTTAA
SEQ2802	TACCAATAGATATTACACCAAAATCTGGTAGTAGAAAAATGAATGGTAGTAAGGTCAC
SEQ2803	TGTCGCTT-ATACTAATCATCAAGAGAA-AAAAACGCAATATAAAGGAC-GTTATTTTAA
	TOTAL THE CATCARGAGAA AAAAACGCAATATAAAGGAC-GTTATTTAA
SEQ2804	TGTCGCTT-ATACTAATCATCAAGAGAA-AAAAACGCAATATAAAGGAC-GTTATTTTAA
SEQ2805	TGTCGCTT-ATACTAATCATCAAGAGAA-AAAAACGCAATATAAAGGAC-GTTATTTTAA
SEQ2806	TOTAL AND COMPANY AND
<del>-</del>	TGTCGCTT-ATACTAATCATCAAGAGAA-AAAAACGCAATATAAAGGAC-GTTATTTTAA
SEQ2807	TGTCGCTT-ATACTAATCATCAAGAGAA-AAAAACGCAATATAAAGGAC-GTTATTTTAA
SEQ2808	TGTCGCTTTATACTAATCATCAAGAGGAGAAAAACGCAATATAAAGGAC-GTTATTTTAA
-	TOTAL THE CALCULATION OF THE CONTROL OF THE CONTROL OF THE CALCULATION
SEQ2809	TGTCGCTT-ATACTAATCATCAAGAGAA-AAAAACGCAATATAAAGGAC-GTTATTTTAA
SEQ2810	TGTCGCTT-ATACTAATCATCAAGAGAA-AAAAACGCAATATAAAGGAC-GTTATTTTAA
SEQ2811	MCMCCCOM AMAGMANAGAROLICAL ABBRICANIA MARAGARO GIRATITAN
25057	TGTCGCTT-ATACTAATCATCAAGAGAA-AAAAACGCAATATAAAGGAC-GTTATTTTAA
SEQ2801	AACTTCTGCGGCAGCTAATCCATTTGAGGTCATGCTAGCTCAAGTTATGGATGAATTG
-	THE TOTAL CONTROL OF THE CANADICATION OF THE C
SEQ2802	ATTTTCTAAATCTAGTGA-CTTTGTATTGTC-TATTGATCCAAATGGCAAGTCTGAATT-
SEQ2803	AACTTCTGTGGCAGCTAATCCATTTGAGGTCATGCTAGCTCAAGTAATGGATGAATTG
SEQ2804	'S CONTROLL OF THE CONTROL OF
	AACTTCTGTGGCAGCTAATCCATTTGAGGTCATGCTAGCTCAAGTAATGGATGAATTG
SEQ2805	AACTTCTGTGGCAGCTAATCCATTTGAGGTCATGCTAAGCTCAAGTAATGGATGAATTG AACTTCTGTGGCAGCTAATCCATTTGAGGTCATGCTAAGCTAATGGATGAATTG
_	AACTTCTGTGGCAGCTAATCCATTTGAGGTCATGCTAGCTCAAGTAATGGATGAATTG
SEQ2806	AACTTCTGTGGCAGCTAATCCATTTGAGGTCATGCTAGCTCAAGTAATGGATGAATTG AACTTCTGTGGCAGCTAATCCATTTGAGGTCATGCTAGCTCAAGTAATGGATGAATTG
SEQ2806 SEQ2807	AACTTCTGTGGCAGCTAATCCATTTGAGGTCATGCTAGCTCAAGTAATGGATGAATTG AACTTCTGTGGCAGCTAATCCATTTGAGGTCATGCTAGCTCAAGTAATGGATGAATTG AACTTCTGTGGCAGCTAATCCATTTGAGGTCATGCTAGCTCAAGTAATGGATGAATTG
SEQ2806 SEQ2807	AACTTCTGTGGCAGCTAATCCATTTGAGGTCATGCTAGCTCAAGTAATGGATGAATTG AACTTCTGTGGCAGCTAATCCATTTGAGGTCATGCTAGCTCAAGTAATGGATGAATTG AACTTCTGTGGCAGCTAATCCATTTGAGGTCATGCTAGCTCAAGTAATGGATGAATTG
SEQ2806 SEQ2807 SEQ2808	AACTTCTGTGGCAGCTAATCCATTTGAGGTCATGCTAGCTCAAGTAATGGATGAATTG AACTTCTGTGGCAGCTAATCCATTTGAGGTCATGCTAGCTCAAGTAATGGATGAATTG AACTTCTGTGGCAGCTAATCCATTTGAGGTCATGCTAGCTCAAGTAATGGATGAATTG AACTTCTGTGGCAGCTAATCCATTTGAGGTCATGCTAGCTCAAGTAATGGATGAATTG
SEQ2806 SEQ2807 SEQ2808 SEQ2809	AACTTCTGTGGCAGCTAATCCATTTGAGGTCATGCTAGCTCAAGTAATGGATGAATTG AACTTCTGTGGCAGCTAATCCATTTGAGGTCATGCTAGCTCAAGTAATGGATGAATTG AACTTCTGTGGCAGCTAATCCATTTGAGGTCATGCTAGCTCAAGTAATGGATGAATTG AACTTCTGTGGCAGCTAATCCATTTGAGGTCATGCTAGCTCAAGTAATGGATGAATTG AACTTCTGTGGCAGCTAATCCATTTGAGGTCATGCTAGCTCAAGTAATGGATGAATTG
SEQ2806 SEQ2807 SEQ2808	AACTTCTGTGGCAGCTAATCCATTTGAGGTCATGCTAGCTCAAGTAATGGATGAATTG AACTTCTGTGGCAGCTAATCCATTTGAGGTCATGCTAGCTCAAGTAATGGATGAATTG AACTTCTGTGGCAGCTAATCCATTTGAGGTCATGCTAGCTCAAGTAATGGATGAATTG AACTTCTGTGGCAGCTAATCCATTTGAGGTCATGCTAGCTCAAGTAATGGATGAATTG AACTTCTGTGGCAGCTAATCCATTTGAGGTCATGCTAGCTCAAGTAATGGATGAATTG
SEQ2806 SEQ2807 SEQ2808 SEQ2809 SEQ2810	AACTTCTGTGGCAGCTAATCCATTTGAGGTCATGCTAGCTCAAGTAATGGATGAATTG AACTTCTGTGGCAGCTAATCCATTTGAGGTCATGCTAGCTCAAGTAATGGATGAATTG AACTTCTGTGGCAGCTAATCCATTTGAGGTCATGCTAGCTCAAGTAATGGATGAATTG AACTTCTGTGGCAGCTAATCCATTTGAGGTCATGCTAGCTCAAGTAATGGATGAATTG AACTTCTGTGGCAGCTAATCCATTTGAGGTCATGCTAGCTCAAGTAATGGATGAATTG AACTTCTGTGGCAGCTAATCCATTTGAGGTCATGCTAGCTCAAGTAATGGATGAATTG
SEQ2806 SEQ2807 SEQ2808 SEQ2809	AACTTCTGTGGCAGCTAATCCATTTGAGGTCATGCTAGCTCAAGTAATGGATGAATTG AACTTCTGTGGCAGCTAATCCATTTGAGGTCATGCTAGCTCAAGTAATGGATGAATTG AACTTCTGTGGCAGCTAATCCATTTGAGGTCATGCTAGCTCAAGTAATGGATGAATTG AACTTCTGTGGCAGCTAATCCATTTGAGGTCATGCTAGCTCAAGTAATGGATGAATTG AACTTCTGTGGCAGCTAATCCATTTGAGGTCATGCTAGCTCAAGTAATGGATGAATTG
SEQ2806 SEQ2807 SEQ2808 SEQ2809 SEQ2810	AACTTCTGTGGCAGCTAATCCATTTGAGGTCATGCTAGCTCAAGTAATGGATGAATTG AACTTCTGTGGCAGCTAATCCATTTGAGGTCATGCTAGCTCAAGTAATGGATGAATTG AACTTCTGTGGCAGCTAATCCATTTGAGGTCATGCTAGCTCAAGTAATGGATGAATTG AACTTCTGTGGCAGCTAATCCATTTGAGGTCATGCTAGCTCAAGTAATGGATGAATTG AACTTCTGTGGCAGCTAATCCATTTGAGGTCATGCTAGCTCAAGTAATGGATGAATTG AACTTCTGTGGCAGCTAATCCATTTGAGGTCATGCTAGCTCAAGTAATGGATGAATTG AACTTCTGTGGCAGCTAATCCATTTGAGGTCATGCTAGCTCAAGTAATGGATGAATTG
SEQ2806 SEQ2807 SEQ2808 SEQ2809 SEQ2810	AACTTCTGTGGCAGCTAATCCATTTGAGGTCATGCTAGCTCAAGTAATGGATGAATTG AACTTCTGTGGCAGCTAATCCATTTGAGGTCATGCTAGCTCAAGTAATGGATGAATTG AACTTCTGTGGCAGCTAATCCATTTGAGGTCATGCTAGCTCAAGTAATGGATGAATTG AACTTCTGTGGCAGCTAATCCATTTGAGGTCATGCTAGCTCAAGTAATGGATGAATTG AACTTCTGTGGCAGCTAATCCATTTGAGGTCATGCTAGCTCAAGTAATGGATGAATTG AACTTCTGTGGCAGCTAATCCATTTGAGGTCATGCTAGCTCAAGTAATGGATGAATTG AACTTCTGTGGCAGCTAATCCATTTGAGGTCATGCTAGCTCAAGTAATGGATGAATTG
SEQ2806 SEQ2807 SEQ2808 SEQ2809 SEQ2810 SEQ2811	AACTTCTGTGGCAGCTAATCCATTTGAGGTCATGCTAGCTCAAGTAATGGAT GAATTG AACTTCTGTGGCAGCTAATCCATTTGAGGTCATGCTAGCTCAAGTAATGGAT GAATTG AACTTCTGTGGCAGCTAATCCATTTGAGGTCATGCTCAAGTAATGGAT GAATTG AACTTCTGTGGCAGCTAATCCATTTGAGGTCATGCTCAAGTAATGGAT GAATTG AACTTCTGTGGCAGCTAATCCATTTGAGGTCATGCTAGCTCAAGTAATGGAT GAATTG AACTTCTGTGGCAGCTAATCCATTTGAGGTCATGCTAGCTCAAGTAATGGAT GAATTG AACTTCTGTGGCAGCTAATCCATTTGAGGTCATGCTAGCTCAAGTAATGGAT GAATTG AACTTCTGTGGCAGCTAAATCCATTTGAGGTCATGCTTAGTTTTTCAAACTCACCA
SEQ2806 SEQ2807 SEQ2808 SEQ2809 SEQ2810 SEQ2811 SEQ2801 SEQ2802	AACTTCTGTGGCAGCTAATCCATTTGAGGTCATGCTAGCTCAAGTAATGGAT GAATTG AACTTCTGTGGCAGCTAATCCATTTGAGGTCATGCTAGCTCAAGTAATGGAT GAATTG AACTTCTGTGGCAGCTAATCCATTTGAGGTCATGCTAGCTCAAGTAATGGAT GAATTG AACTTCTGTGGCAGCTAATCCATTTGAGGTCATGCTAGCTCAAGTAATGGAT GAATTG AACTTCTGTGGCAGCTAATCCATTTGAGGTCATGCTAGCTCAAGTAATGGAT GAATTG AACTTCTGTGGCAGCTAATCCATTTGAGGTCATGCTAGCTCAAGTAATGGAT GAATTG AACTTCTGTGGCAGCTAATCCATTTGAGGTCATGCTCAAGTAATGGAT GAATTG ACACATTATGAGACAGCTAAATATGGTTGGCAACATTTGATTAGTTTTTCAAACTCACCA ATTTGTC-CAAGAGCGCTATA-ATGCCT TAAAAGCGAACTATCTTCGACAGCTTAACG
SEQ2806 SEQ2807 SEQ2808 SEQ2809 SEQ2810 SEQ2811	AACTTCTGTGGCAGCTAATCCATTTGAGGTCATGCTAGCTCAAGTAATGGAT GAATTG AACTTCTGTGGCAGCTAATCCATTTGAGGTCATGCTAGCTCAAGTAATGGAT GAATTG AACTTCTGTGGCAGCTAATCCATTTGAGGTCATGCTAGCTCAAGTAATGGAT GAATTG AACTTCTGTGGCAGCTAATCCATTTGAGGTCATGCTAGCTCAAGTAATGGAT GAATTG AACTTCTGTGGCAGCTAATCCATTTGAGGTCATGCTAGCTCAAGTAATGGAT GAATTG AACTTCTGTGGCAGCTAATCCATTTGAGGTCATGCTAGCTCAAGTAATGGAT GAATTG AACTTCTGTGGCAGCTAATCCATTTGAGGTCATGCTCAAGTAATGGAT GAATTG ACACATTATGAGACAGCTAAATATGGTTGGCAACATTTGATTAGTTTTTCAAACTCACCA ATTTGTC-CAAGAGCGCTATA-ATGCCT TAAAAGCGAACTATCTTCGACAGCTTAACG
SEQ2806 SEQ2807 SEQ2808 SEQ2809 SEQ2810 SEQ2811 SEQ2801 SEQ2802 SEQ2803	AACTTCTGTGGCAGCTAATCCATTTGAGGTCATGCTAGCTCAAGTAATGGAT GAATTG AACTTCTGTGGCAGCTAATCCATTTGAGGTCATGCTAGCTCAAGTAATGGAT GAATTG AACTTCTGTGGCAGCTAATCCATTTGAGGTCATGCTAGCTCAAGTAATGGAT GAATTG AACTTCTGTGGCAGCTAATCCATTTGAGGTCATGCTAGCTCAAGTAATGGAT GAATTG AACTTCTGTGGCAGCTAATCCATTTGAGGTCATGCTAGCTCAAGTAATGGAT GAATTG AACTTCTGTGGCAGCTAATCCATTTGAGGTCATGCTCAAGTAATGGAT GAATTG AACTTCTGTGGCAGCTAATCCATTTGAGGTCATGCTCAAGTAATGGAT GAATTG ACACATTATGAGACAGCTAAATATGGTTGGCAACATTTGATTAGTTTTTCAAACTCACCA ATTTGTC - CAAGAGCGCTATA - ATGCCT TAAAAGCGAACTATCTTCGACAGCTTAACG ACACATTATGAGACAGCTAAATATGGTTGGCAACATTTGATTAGTTTTTCAAACTCACCA
SEQ2806 SEQ2807 SEQ2808 SEQ2809 SEQ2810 SEQ2811 SEQ2801 SEQ2802	AACTTCTGTGGCAGCTAATCCATTTGAGGTCATGCTAGCTCAAGTAATGGAT GAATTG AACTTCTGTGGCAGCTAATCCATTTGAGGTCATGCTAGCTCAAGTAATGGAT GAATTG AACTTCTGTGGCAGCTAATCCATTTGAGGTCATGCTAGCTCAAGTAATGGAT GAATTG AACTTCTGTGGCAGCTAATCCATTTGAGGTCATGCTAGCTCAAGTAATGGAT GAATTG AACTTCTGTGGCAGCTAATCCATTTGAGGTCATGCTAGCTCAAGTAATGGAT GAATTG AACTTCTGTGGCAGCTAATCCATTTGAGGTCATGCTAGCTCAAGTAATGGAT GAATTG AACTTCTGTGGCAGCTAATCCATTTGAGGTCATGCTCAAGTAATGGAT GAATTG ACACATTATGAGACAGCTAAATATGGTTGGCAACATTTGATTAGTTTTTCAAACTCACCA ATTTGTC-CAAGAGCGCTATA-ATGCCT TAAAAGCGAACTATCTTCGACAGCTTAACG



SEQ2805	ACACATTATGAGACAGCTAAATATGGTTGGCAACATTTGATTAGTTTTTCAAACTCACCA
SEQ2806	ACACATTATGAGACAGCTAAATATGGTTGGCAACATTTGATTAGTTTTTCAAACTCACCA
	ACACATTA GAGACAGCTAAATA GGTTGGCAACATTTGATTAGTTTTCAAACTCACCA
SEQ2807	ACACATTATGAGACAGCTAAATATGGTTGGCAACATTTGATTAGTTTTTCAAACTCACCA
SEQ2808	ACACATTATGAGACAGCTAAATATGGTTGGCAACATTTGATTAGTTTTTCAAACTCACCA
SEQ2809	ACACATTATGAGACAGCTAAATATGGTTGGCAACATTTGATTAGTTTTTCAAACTCACCA
SEQ2810	ACACATTATGAGACAGCTAAATATGGTTGGCAACATTTGATTAGTTTTTCAAACTCACCA
SEQ2811	ACACATTATGAGACAGCTAAATATGGTTGGCAACATTTGATTAGTTTTTCAAACTCACCA
SEQ2801	CAACAGACCCTTTTCGTTATCGAAAACCATTTG-AGGCACAGGCTCCTAAATA
SEQ2802	TAAAGATTTTTATGCTTTCCCACCAAAGAAGAACAGTAGTAATTTTGAGCAGATCAATA
SEQ2803	CAACAGACCCTTTTCATTATCGAAAACCATTTG-AGGCACAGGCTCCTAAATA
SEQ2804	CAACAGACCCTTTTCATTATCGAAAACCATTTG-AGGCACAGGCTCCTAAATA
SEQ2805	CAACAGACCCTTTTCATTATCGAAAACCATTTG-AGGCACAGGCTCCTAAATA
SEQ2806	CAACAGACCCTTTTCATTATCGAAAACCATTTG-AGGCACAGGCTCCTAAATA
SEQ2807	CAACAGACCCTTTTCATTATCGAAAACCATTTG-AGGCACAGGCTCCTAAATA
SEQ2808	CAACAGACCCTTTTCATTATCGAAAACCATTTG-AGGCACAGGCTCCTAAATA
SEQ2809	CAACAGACCCTTTTCATTATCGAAAACCATTTG-AGGCACAGGCTCCTAAATA
SEQ2810	CAACAGACCCTTTTCATTATCGAAAACCATTTG-AGGCACAGGCTCCTAAATA
SEQ2811	CAACAGACCCTTTTCATTATCGAAAACCATTTG-AGGCACAGGCTCCTAAATA
•	
SEQ2801	G-TACAACTAAATGTAGAAAATATTCAAGCTAATTCGAATGTTAAAGCAGGTATTT
SEQ2802	GGTATTGAGAAATACAAAGATTGTTGAAGACATGGAAAAAGTAAAAGCAACAGAGAGGT
SEQ2803	G-TACAACTAAATGTAGAAAATATTCAAGCTAATTCAAATGTTAAAGCAGGTATGT
SEQ2804	G-TACAACTAAATGTAGAAAATATTCAAGCTAATTCAAATGTTAAAGCAGGTATGT
SEQ2805	G-TACAACTAAATGTAGAAAATATTCAAGCTAATTCAAATGTTAAAGCAGGTATGT
SEQ2806	G-TACAACTAAATGTAGAAAATATTCAAGCTAATTCAAATGTTAAAGCAGGTATGT
SEQ2807	G-TACAACTAAATGTAGAAAATATTCAAGCTAATTCAAATGTTAAAGCAGGTATGT
SEQ2808	G-TACAACTAAATGTAGAAAATATTCAAGCTAATTCGAATGTTAAAGCAGGTATGT
SEQ2809	G-TACAACTAAATGTAGAAAATATTCAAGCTAATTCGAATGTTAAAGCAGGTATGT
SEQ2810	G-TACAACTAAATGTAGAAAATATTCAAGCTAATTCAAATGTTAAAGCAGGTATGT
SEQ2811	G-TACAACTAAATGTAGAAAATATTCAAGCTAATTCAAATGTTAAAGCAGGTATGT
SEQ2801	TTGCAGCATATAAAGCTATTGATTTCCATCCTCGATACAAGGATTATCTATTATTTGATA
SEQ2802	TCTTACCAACTCATCCTACTGGTCTTCTCAAAACAGGAACAATTGAT-AGGCACCA
SEQ2803	TTGCAGCATATAAAGCTATTGATTTCCATCCTCGATACAAGGATTATCTATTATTTGATA
'SEQ2804	TTGCAGCATATAAAGCTATTGATTCCATCCTCGATACAAGGATTATCTATTATTTGATA
SEQ2805	TTGCAGCATATAAAGCTATTGATTTCCATCCTCGATACAAGGATTATCTATTATTTGATA
SEQ2806	TTGCAGCATATAAAGCTATTGATTTCCATCCTCGATACAAGGATTATCTATTATTTGATA
SEQ2807	TTGCAGCATATAAAGCTATTGATTTCCATCCTCGATACAAGGATTATCTATTATTTGATA
SEQ2808	TTGCAGCATATAAAGCTATTGATTTCCATCCTCGATACAAGGATTATCTATTATTTGATA
SEQ2809	TTGCAGCATATAAAGCTATTGATTTCCATCCTCGATACAAGGATTATCTATTATTTGATA
SEQ2810	TTGCAGCATATAAAGCTATTGATTTCCATCCTCGATACAAGGATTATCTATTATTTGATA
SEQ2811	TTGCAGCATATAAAGCTATTGATTTCCATCCTCGATACAAGGATTATCTATTATTTGATA
SEQ2801	AAGAGAATATCAGTAAAGAAGATAGACAAAAGATT-AAAGAACTTTCTTTGTCACAGGGA
SEQ2802	AAAAACATTTGATTCACAAACAGATATTTCGTTTGGAAAGGACTTTATAGAGGTCAGAAT
SEQ2803	AAGAGAATATCAGTAAAGAAGATAGACAAAAGATT-AAAGAACTTTCTTTGTCACAGGGA
SEQ2804	AAGAGAATATCAGTAAAGAAGATAGACAAAAGATT-AAAGAACTTTCTTTGTCACAGGGA
SEQ2805	AAGAGAATATCAGTAAAGAAGATAGACAAAAGATT-AAAGAACTTTCTTTGTCACAGGGA
SEQ2806	AAGAGAATATCAGTAAAGAAGATAGACAAAAGATT-AAAGAACTTTCTTTGTCACAGGGA
SEQ2807	AAGAGAATATCAGTAAAGAAGATAGACAAAAGATT-AAAGAACTTTCTTTGTCACAGGGA
SEQ2808	AAGAGAATATCAGTAAAGAAGATAGACAAAAGATT-AAAGAACTTTCTTTGTCACAGGGA
SEQ2809	AAGAGAATATCAGTAAAGAAGATAGACAAAAGATT-AAAGAACTTTCTTTGTCACAGGGA
SEQ2810	AAGAGAATATCAGTAAAGAAGATAGACAAAAGATT-AAAGAACTTTCTTTGTCACAGGGA
SEQ2811	AAGAGAATATCAGTAAAGAAGATAGACAAAAGATT-AAAGAACTTTCTTTGTCACAGGGA
SEQ2801	TACGTTA-AACTGCTAAATGCTTATCACAAAATCCCTGTTCTAGTCACGGGTTATGGCTA
SEQ2802	TCCGTGGCAGTTGTTGAATTTTTCTGATCCATCATCTCAAAAAATTCACGATGATTA
SEQ2803	TACGTTA-AACTGCTAAATGCTTATCACAAAATCCCTGTTCTAGTCACGGGTTATGGCTA
SEQ2804	TACGTTA-AACTGCTAAATGCTTATCACAAAATCCCTGTTCTAGTCACGGGTTATGGCTA
SEQ2805	TACGTTA-AACTGCTAAATGCTTATCACAAAATCCCTGTTCTAGTCACGGGTTATGGCTA
SEQ2806	TACGTTA-AACTGCTAAATGCTTATCACAAAATCCCTGTTCTAGTCACGGGTTATGGCTA
SEQ2807	TACGTTA-AACTGCTAAATGCTTATCACAAAATCCCTGTTCTAGTCACGGGTTATGGCTA
SEQ2808	TACGTTA-AACTGCTAAATGCTTATCACAAAATCCCTGTTCTAGTCACGGGTTATGGCTA
SEQ2809	TACGTTA-AACTGCTAAATGCTTATCACAAAATCCCTGTTCTAGTCACGGGTTATGGCTA
SEQ2810	TACGTTA-AACTGCTAAATGCTTATCACAAAATCCCTGTTCTAGTCACGGGTTATGGCTA TACGTTA-AACTGCTAAATGCTTATCACAAAATCCCTGTTCTAGTCACGGGTTATGGCTA
SEQ2811	TACGTTA-AACTGCTAAATGCTTATCACAAAATCCCTGTTCTAGTCACGGGTTATGGCTA TACGTTA-AACTGCTAAATGCTTATCACAAAATCCCTGTTCTAGTCACGGGTTATGGCTA



# Tabl 28: Comparative Sequences relating t SAG1552 (conserved hypothetical pr tein)

_SEQ2801	TCGACAGCGAGAGGTATTGCCCAAAAAGAAATTGATAAACGTCCTCTGCCGATTAATGA
	TTTAAACATTATGGTGTGAAGGAGTTAGAAATTGA-GAGCATTGCTTTAGGATTAGGTG
SEQ2802	II IMAACAI IAIGGIGIGAAGGAGI IAGAAAI IGA—GAGCATIGCITTAGGATIAGGIG
SEQ2803	TCGACAGCGAGAGGTATTGCCCAAAAAGAAATTGATAAACGTCCTCTGCCGATTAATGA
-	TCGACAGCGAGAGGTATTGCCCAAAAAGAAATTGATAAACGTCCTCTGCCGATTAATGA
SEQ2804	TCGACAGCGAGAGGTATTGCCCCAAAAAGAAATTGATAAACGTCCTCTGCCGATTAATGA
SEQ2805	TCGACAGCGAGAGGTATTGCCCAAAAAGAAATTGATAAACGTCCTCTGCCGATTAATGA
SEQ2806	TCGACAGCGAGAGGTATTGCCCAAAAAGAAATTGATAAACGTCCTCTGCCGATTAATGA
	TCGACAGCGAGAGGTATTGCCCAAAAAGAAATTGATAAACGTCCTCCCGATTAATGA
SEQ2807	TCGACAGCGAGAGGTATTGCCCAAAAAGAAATTGATAAACGTCCTCTGCCGATTAATGA
SEQ2808	TCGACAGCGAGAGGTATTGCCCAAAAAGAAATTGATAAACGTCCTCTGCCGATTAATGA
	TCGACAGCGAGAGGTATTGCCCCAAAAAGAAATTGATAAACGTCCTCTGCCGATTAATGA
SEQ2809	TCGACAGCGAGAGGTATTGCCCAAAAAGAAATTGATAAACGTCCTCTGCCGATTAATGA
-	MOCA CA COCA CA COMA MACACANA A A A A A A A A A A A A A A A A
SEQ2810	TCGACAGCGAGAGGTATTGCCCAAAAAGAAATTGATAAACGTCCTCTGCCGATTAATGA
SEQ2811	TCGACAGCGAGAGGTATTGCCCAAAAAGAAATTGATAAACGTCCTCTGCCGATTAATGA
SEQ2801	AAAGAACAAGGTCAGCGTTTACTAGAAGATTATGAATCTTTTATATCATCCGGTAGTTT
SEQ2802	TAATAGCAAAGAAAACACTGATAAAGATGGCAGATTATCGTTTGAAAAATT
SEQ2803	AAAGAACAAGGTCAGCGTTTACTAGAAGATTATGAATCTTTTATATCATCCGGTAGTTT
SEQ2804	AAAGAACAAGGTCAGCGTTTACTAGAAGATTATGAATCTTTTATATCATCCGGTAGTTT
SEQ2805	AAAGAACAAGGTCAGCGTTTACTAGAAGATTATGAATCTTTTATATCATCCGGTAGTTT
SEQ2806	AAAGAACAAGGTCAGCGTTTACTAGAAGATTATGAATCTTTTATATCATCCGGTAGTTT
SEQ2807	AAAGAACAAGGTCAGCGTTTACTAGAAGATTATGAATCTTTTATATCATCCGGTAGTTT
SEQ2808	
	AAAGAACAAGGTCAGCGTTTACTAGAAGATTATGAATCTTTTATATCATCCGGTAGTTT
SEQ2809	AAAGAACAAGGTCAGCGTTTACTAGAAGATTATGAATCTTTTATATCATCCGGTAGTTT
SEQ2810	AAAGAACAAGGTCAGCGTTTACTAGAAGATTATGAATCTTTTATATCATCCGGTAGTTT
SEQ2811	AAAGAACAAGGTCAGCGTTTACTAGAAGATTATGAATCTTTTATATCATCCGGTAGTTT
SEQ2801	GGAGCGACTATCAATGCATGGCAAGACGATTGGAATGCAAGGGCGTGGAATACATCCTT
SEQ2802	GGAGAGACCCGATACCAAAACCTTTTTAAAAGACTCCTATTATAGTATT
SEQ2803	GGAGCGACTATCAATGCATGGCAAGACGATTGGAATGCAAGGGCGTGGAATACATCTTT
-	
SEQ2804	GGAGCGACTATCAATGCATGGCAAGACGATTGGAATGCAAGGGCGTGGAATACATCTTT
SEQ2805	GGAGCGACTATCAATGCATGGCAAGACGATTGGAATGCAAGGGCGTGGAATACATCTTT
SE02806	
	GGAGCGACTATCAATGCATGGCAAGACGATTGGAATGCAAGGGCGTGGAATACATCTTT
SEQ2807	GGAGCGACTATCAATGCATGGCAAGACGATTGGAATGCAAGGGCGTGGAATACATCTTT
SEQ2808	
	GGAGCGACTATCAATGCATGGCAAGACGATTGGAATGCAAGGGTGTGGAATACATCCTT
SEQ2809	GGAGCGACTATCAATGCATGGCAAGACGATTGGAATGCAAGGGTGTGGAATACATCCTT
SEQ2810	GGAGCGACTATCAATGCATGGCAAGACGATTGGAATGCAAGGGCGTGGAATACATCTTT
SEQ2811	GGAGCGACTATCAATGCATGGCAAGACGATTGGAATGCAAGGGCGTGGAATACATCTTT
4500001	
SEQ2801	GCCACAAATAAACATAGTCAATTCCTATGGGGGGATGCACAAGTATTTAATCAAGGTTA
SEQ2802	AAGAAAGAA
SEQ2803	
250500	COOLOR
	GCCACAAATAAACATAGTCAATTCCTATGGGGGGATGCACAAGTATTTAATCAAGGTTA
SEQ2804	
	GCCACAAATAAACATAGTCAATTCCTATGGGGGGATGCACAAGTATTTAATCAAGGTTA
SEQ2805	GCCACAAATAAACATAGTCAATTCCTATGGGGGGATGCACAAGTATTTAATCAAGGTTA GCCACAAATAAACATAGTCAATTCCTATGGGGGGATGCACAAGTATTTAATCAAGGTTA
	GCCACAAATAAACATAGTCAATTCCTATGGGGGGATGCACAAGTATTTAATCAAGGTTA GCCACAAATAAACATAGTCAATTCCTATGGGGGGATGCACAAGTATTTAATCAAGGTTA
SEQ2805 SEQ2806	GCCACAAATAAACATAGTCAATTCCTATGGGGGGATGCACAAGTATTTAATCAAGGTTA GCCACAAATAAACATAGTCAATTCCTATGGGGGGATGCACAAGTATTTAATCAAGGTTA GCCACAAATAAACATAATCAATTCCTATGGGGGGATGCACAAGTATTTAATCAAGGTTA
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SEQ2805 SEQ2806 SEQ2807 SEQ2808 SEQ2809	GCCACAAATAAACATAGTCAATTCCTATGGGGGGATGCACAAGTATTTAATCAAGGTTA GCCACAAATAAACATAGTCAATTCCTATGGGGGGATGCACAAGTATTTAATCAAGGTTA GCCACAAATAAACATAATCAATTCCTATGGGGGGATGCACAAGTATTTAATCAAGGTTA GCCACAAATAAACATAGTCAATTCCTATGGGGGGATGCACAAGTATTTAATCAAGGTTA GCCACAAATAAACATAGTCAATTCCTATGGGGGGATGCACAAGTATTTAATCAAGGTTA GCCACAAATAAACATAGTCAATTCCTATGGGGGGATGCACAAGTATTTAATCAAGGTTA
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SEQ2805 SEQ2806 SEQ2807 SEQ2809 SEQ2810 SEQ2811 SEQ2801 SEQ2802 SEQ2803 SEQ2804 SEQ2805 SEQ2806 SEQ2807 SEQ2808 SEQ2809 SEQ2810 SEQ2811 SEQ2801 SEQ2801 SEQ2801 SEQ2801	GCCACAAATAAACATAGTCAATTCCTATGGGGGGATGCACAAGTATTTAATCAAGGTTA GGTTTATTAGGCTTTAAAAACGCAAAACATCATTATCAAGTTGATGGTAAAAGAGCAA GGTTTATTAGGCTTTAAAAACGCAAAACATCATTATCAAGTTGATGGTAAAAGAGCAA GGTTTATTAGGCTTTAAAAACGCAAAACATCATTATCAAGTTGATGGTAAAAGAGCCAA GGTTTATTAGGCTTTAAAAACGCAAAACATCATTATCAAGTTGATGGTAAAAGAGGCAA GGTTTATTAGGCTTTAAAAACGCAAAACATCATTATCAAGTTGATGGTAAAAGAGGCAA GGTTTATTAGGCTTTAAAAACGCAAAACATCATTATCAAGTTGATGGTAAAAGAGCCAA GGTTTATTAGGCTTTAAAAACGCAAAACATCATTATCAAGTTGATGGTAAAAGAGCCAA GGTTTATTAGGCTTTAAAAACGCAAAACATCATTATCAAGTTGATGGTAAAAGAGCCAA GGTTTATTAGGCTTTAAAAACGCAAAACATCATTATCAAGTTGATGGTAAAAGAGCCAA GGAGAGTGGAAACATCCTCTG
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SEQ2805 SEQ2806 SEQ2807 SEQ2809 SEQ2810 SEQ2811 SEQ2801 SEQ2802 SEQ2803 SEQ2804 SEQ2805 SEQ2806 SEQ2807 SEQ2808 SEQ2809 SEQ2810 SEQ2811 SEQ2801 SEQ2801 SEQ2802 SEQ2803	GCCACAAATAAACATAGTCAATTCCTATGGGGGGATGCACAAGTATTTAATCAAGGTTA GCCACAAATAAACATAGTCAATTCCTATGGGGGGATGCACAAGTATTTAATCAAGGTTA GCCACAAATAAACATAATCCAATTCCTATGGGGGGATGCACAAGTATTTAATCAAGGTTA GCCACAAATAAACATAGTCAATTCCTATGGGGGGATGCACAAGTATTTAATCAAGGTTA GCCACAAATAAACATAGTCAATTCCTATGGGGGGATGCACAAGTATTTAATCAAGGTTA GCCACAAATAAACATAGTCAATTCCTATGGGGGGATGCACAAGTATTTAATCAAGGTTA GCCACAAATAAACATAGTCAATTCCTATGGGGGGATGCACAAGTATTTAATCAAGGTTA GCCACAAATAAACATAGTCAATTCCTATGGGGGGATGCACAAGTATTTAATCAAGGTTA GCCACAAATAAACATAGTCAATTCCTATGGGGGGATGCACAAGTATTTAATCAAGGTTA GCCACAAATAAACATAGTCAATTCCTATGGGGGGATGCACAAGTATTTAATCAAGGTTA GCTTTATTAGGCTTTAAAAACGCAAAACATCATTATCAAGTTGATGGTAAAAGAGGCAA GGTTTATTAGGCTTTAAAAACGCAAAACATCATTATCAAGTTGATGGTAAAAGAGGCAA GGTTTATTAGGCTTTAAAAACGCAAAACATCATTATCAAGTTGATGGTAAAAAGAGGCAA GGAGGTGGAAACATCCTCTGG
SEQ2805 SEQ2806 SEQ2807 SEQ2809 SEQ2810 SEQ2811 SEQ2801 SEQ2802 SEQ2803 SEQ2804 SEQ2805 SEQ2806 SEQ2807 SEQ2808 SEQ2809 SEQ2810 SEQ2811 SEQ2801 SEQ2802 SEQ2803 SEQ2803 SEQ2804	GCCACAAATAAACATAGTCAATTCCTATGGGGGGATGCACAAGTATTTAATCAAGGTTA GCCACAAATAAACATAGTCAATTCCTATGGGGGGATGCACAAGTATTTAATCAAGGTTA GCCACAAAATAAACATAATCCAATTCCTATGGGGGGATGCACAAGTATTTAATCAAGGTTA GCCACAAATAAACATAGTCAATTCCTATGGGGGGATGCACAAGTATTTAATCAAGGTTA GCCACAAAATAAACATAGTCAATTCCTATGGGGGGATGCACAAGTATTTAATCAAGGTTA GCCACAAAATAAACATAGTCAATTCCTATGGGGGGATGCACAAGTATTTAATCAAGGTTA GCCACAAATAAACATAGTCAATTCCTATGGGGGGATGCACAAGTATTTAATCAAGGTTA GCCACAAATAAACATAGTCAATTCCTATGGGGGGATGCACAAGTATTTAATCAAGGTTA GCCACAAATAAACATAGTCAATTCCTATGGGGGGATGCACAAGTATTTAATCAAGGTTA GCCACAAATAAACATAGTCAATTCCTATGGGGGGATGCACAAGTATTTAATCAAGGTTA GCTTTATTAGGCTTTAAAAACGCAAAACATCATTATCAAGTTGATGGTAAAAGAGGCAA GGTTTATTAGGCTTTAAAAACGCAAAACATCATTATCAAGTTGATGGTAAAAGAGGCAA GGAGAGGGAAACATCCTCTGGTGCAACAGGAAGATGACTTATATGCTAGCAG GGAGAGTGGAAACATCCTCTCGATGACTAGTGCAACAGGAAGATGACTTATATGCTAGCAG GGAGAGTGGAAACATCCTCTCGATGACTAGTGCAACAGGAAGATGACTTATATGCTAGCAG GGAGAGTGGAAACATCCTCTCTAGTGACTAGTGCAACAGGAAGATGACTTATATGCTAGCAG GGAGAGTGGAAACATCCTCTCTAGTGACTAGTGCAACAGGAAGATGACTTATATTATGCTAGCAG
SEQ2805 SEQ2806 SEQ2807 SEQ2809 SEQ2810 SEQ2811 SEQ2801 SEQ2802 SEQ2803 SEQ2804 SEQ2805 SEQ2806 SEQ2807 SEQ2808 SEQ2809 SEQ2810 SEQ2811 SEQ2801 SEQ2801 SEQ2802 SEQ2803	GCCACAAATAAACATAGTCAATTCCTATGGGGGGATGCACAAGTATTTAATCAAGGTTA GCCACAAATAAACATAGTCAATTCCTATGGGGGGATGCACAAGTATTTAATCAAGGTTA GCCACAAAATAAACATAATCCAATTCCTATGGGGGGATGCACAAGTATTTAATCAAGGTTA GCCACAAATAAACATAGTCAATTCCTATGGGGGGATGCACAAGTATTTAATCAAGGTTA GCCACAAAATAAACATAGTCAATTCCTATGGGGGGATGCACAAGTATTTAATCAAGGTTA GCCACAAAATAAACATAGTCAATTCCTATGGGGGGATGCACAAGTATTTAATCAAGGTTA GCCACAAATAAACATAGTCAATTCCTATGGGGGGATGCACAAGTATTTAATCAAGGTTA GCCACAAATAAACATAGTCAATTCCTATGGGGGGATGCACAAGTATTTAATCAAGGTTA GCCACAAATAAACATAGTCAATTCCTATGGGGGGATGCACAAGTATTTAATCAAGGTTA GCCACAAATAAACATAGTCAATTCCTATGGGGGGATGCACAAGTATTTAATCAAGGTTA GCTTTATTAGGCTTTAAAAACGCAAAACATCATTATCAAGTTGATGGTAAAAGAGGCAA GGTTTATTAGGCTTTAAAAACGCAAAACATCATTATCAAGTTGATGGTAAAAGAGGCAA GGAGAGGGAAACATCCTCTGGTGCAACAGGAAGATGACTTATATGCTAGCAG GGAGAGTGGAAACATCCTCTCGATGACTAGTGCAACAGGAAGATGACTTATATGCTAGCAG GGAGAGTGGAAACATCCTCTCGATGACTAGTGCAACAGGAAGATGACTTATATGCTAGCAG GGAGAGTGGAAACATCCTCTCTAGTGACTAGTGCAACAGGAAGATGACTTATATGCTAGCAG GGAGAGTGGAAACATCCTCTCTAGTGACTAGTGCAACAGGAAGATGACTTATATTATGCTAGCAG
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SEQ2805 SEQ2806 SEQ2807 SEQ2809 SEQ2810 SEQ2811 SEQ2801 SEQ2803 SEQ2804 SEQ2805 SEQ2806 SEQ2806 SEQ2809 SEQ2810 SEQ2811 SEQ2801 SEQ2801 SEQ2802 SEQ2803 SEQ2804 SEQ2804 SEQ2805 SEQ2806 SEQ2806	GCCACAAATAAACATAGTCAATTCCTATGGGGGGATGCACAAGTATTTAATCAAGGTTA GCTTTATTAGGCTTTAAAAACGCAAAACATCATTATCAAGTTGATGGTAAAAGAGGCAA GGTTTATTAGGCTTTAAAAACGCAAAACATCATTATCAAGTTGATGGTAAAAGAGGCAA GGTTAATTAGGCTTTAAAAACGCAAAACATCATTATCAAGTTGATGGTAAAAGAGGCAA GGAGATGGAAACATCCTCTGATGACTAGTGCAACAGGAGATGACTTATATGCTAGCAG GGAGATGGAAACATCCTCTGATGACTAGTGCAACAGGAGATGACTTATATGCTAGCAG GGAGATGGAAACATCCTCTGATGACTAGTGCAACAGGAGATGACTTATATGCTAGCAG GGAGATGGAAACATCCTCTGATGACTAGTGCAACAGGAGATGACTTATATGCTAGCAG GGAGATGGAAACATCCTCTGATGACTAGTGCAACAGGAGATGACTTATATGCTAGCAG GGAGATGGAAACATCCTCTTATTAGCTAGCTAGTGCAACAGGAGATGACTTATATGCTAGCAG GGAGATGGAAACATCCTCTGATGACTAGTGCAACAGGAGATGACTTATATGCTAGCAG GGAGATGGAAACATCCTCTGATGACTAGTGCAACAGGAGATGACTTATATATGCTAGCAG GGAGAGT
SEQ2805 SEQ2806 SEQ2807 SEQ2809 SEQ2810 SEQ2811 SEQ2801 SEQ2803 SEQ2803 SEQ2804 SEQ2805 SEQ2806 SEQ2809 SEQ2810 SEQ2811 SEQ2801 SEQ2802 SEQ2803 SEQ2804 SEQ2804 SEQ2804 SEQ2804 SEQ2804	GCCACAAATAAACATAGTCAATTCCTATGGGGGGATGCACAAGTATTTAATCAAGGTTA GCTTTATTAGGCTTTAAAAACGCAAAACATCATTATCAAGTTGATGGTAAAAGAGGCAA GGTTTATTAGGCTTTAAAAACGCAAAACATCATTATCAAGTTGATGGTAAAAGAGGCAA GGTTAATTAGGCTTTAAAAACGCAAAACATCATTATCAAGTTGATGGTAAAAGAGGCAA GGAGATGGAAACATCCTCTGATGACTAGTGCAACAGGAGATGACTTATATGCTAGCAG GGAGATGGAAACATCCTCTGATGACTAGTGCAACAGGAGATGACTTATATGCTAGCAG GGAGATGGAAACATCCTCTGATGACTAGTGCAACAGGAGATGACTTATATGCTAGCAG GGAGATGGAAACATCCTCTGATGACTAGTGCAACAGGAGATGACTTATATGCTAGCAG GGAGATGGAAACATCCTCTGATGACTAGTGCAACAGGAGATGACTTATATGCTAGCAG GGAGATGGAAACATCCTCTTATTAGCTAGCTAGTGCAACAGGAGATGACTTATATGCTAGCAG GGAGATGGAAACATCCTCTGATGACTAGTGCAACAGGAGATGACTTATATGCTAGCAG GGAGATGGAAACATCCTCTGATGACTAGTGCAACAGGAGATGACTTATATATGCTAGCAG GGAGAGT
SEQ2805 SEQ2806 SEQ2807 SEQ2809 SEQ2810 SEQ2811 SEQ2801 SEQ2803 SEQ2804 SEQ2805 SEQ2806 SEQ2806 SEQ2809 SEQ2810 SEQ2811 SEQ2801 SEQ2801 SEQ2802 SEQ2803 SEQ2804 SEQ2804 SEQ2805 SEQ2806 SEQ2806	GCCACAAATAAACATAGTCAATTCCTATGGGGGGATGCACAAGTATTTAATCAAGGTTA GCTATATTAGGCTTTAAAAACGCAAAACATCATTATCAAGTTGATGGTAAAAGAGGCAA GGTTTATTAGGCTTTAAAAACGCAAAACATCATTATCAAGTTGATGGTAAAAGAGGCAA GGTTTATTAGGCTTTAAAAACGCAAAACATCATTATCAAGTTGATGGTAAAAGAGGCAA GGTTTATTAGGCTTTAAAAACGCAAAACATCATTATCAAGTTGATGGTAAAAGAGGCAA GGTTTATTAGGCTTTAAAAACGCAAAACATCATTATCAAGTTGATGGTAAAAGAGGCAA GGTTTATTAGGCTTTAAAAACGCAAAACATCATTATCAAGTTGATGGTAAAAGAGGCAA GGTTTATTAGGCTTTAAAAACGCAAAACATCATTATCAAGTTGATGGTAAAAGAGGCAA GGTTTATTAGGCTTTAAAAACGCAAAACATCATTATCAGGTTGATGGTAAAAGAGGCAA GGTTTATTAGGCTTTAAAAACGCAAAACATCATTATCAGGTTGATGGTAAAAGAGGCAA GGTTTATTAGGCTTTAAAAACGCAAAACATCATTATCAGGTTGATGGTAAAAGAGGCAA GGTTTATTAGGCTTTAAAAACGCAAAACATCATTATCAGGTTGATGGTAAAAGAGGCAA GGTTTATTAGGCTTTAAAAACGCAAAACATCATTATCAGGTTGATGGTAAAAGAGGCAA GGTTTATTAGGCTTTAAAAACGCAAAACATCATTATCAGGTTGATGGTAAAAGAGGCAA GGTTTATTAGGCTTTAAAAACGCAAAACATCATTATCAGGTTGATGGTAAAAGAGGCAA GGTTTATTAGGCTTTAAAAACGCAAAACATCATTATCAGGTTGATGGTAAAAGAGGCAA GGTTTATTAGGCTTTAAAAACGCAAAACATCATTATCAGGTTGATGGTAAAAGAGGCAA GGTTTATTAGGCTTTAAAAACGCAAAACATCATTATCAAGTTGATGGTAAAAGAGGCAA GGAGAGTGGAAACATCCTCTGATGACTAGTGCAACAGGAGATGACTTATATGCTAGCAG GGAGAGTGGAAACATCCTCTGATGACTAGTGCAACAGGAGATGACTTATATGCTAGCAG GGAGAGTGGAAACATCCTCTGATGACTAGTGCAACAGGAGATGACTTATATGCTAGCAG GGAGAGTGGAAACATCCTCTGATGACTAGTGCAACAGGAGATGACTTATATTATTATGCTAGCAG GGAGAGTGGAAACATCCTCTGATGACTAGTGCAACAGGAGATGACTTATATATGCTAGCAG GGAGAGTGGAAACATCCTCTGATGACTAGTGCAACAGGAGATGACTTATATATGCTAGCAG

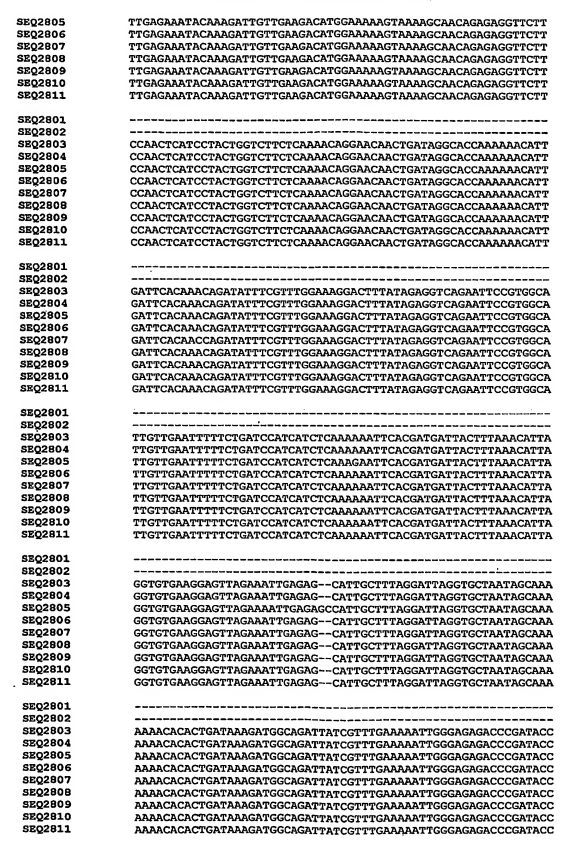


# 604062 Table 28: Comparative Sequences relating to SAG1552 (conserved hypothetical protein)

	•
SEQ2809	GAAGAGTGGAAACATCCTCTGATGACTAGTGCAACAGGAGATGACTTATATGCTAGCAG
SEQ2810	GGAGAGTGGAAACATCCTCTGATGACTAGTGCAACAGGAGATGACTTATATGCTAGCAG
<del></del>	
SEQ2811	GGAGAGTGGAAACATCCTCTGATGACTAGTGCAACAGGAGATGACTTATATGCTAGCAG
SEQ2801	
SEQ2802	
SEQ2803	GATGAAAGCTATCTCTACCTTGCGATTAAAACAAAACCTGAAAAACTAAAAGAAAAACG
SEQ2804	GATGAAAGCTATCTCTACCTTGCGATTAAAACAAAACCTGAAAAACTAAAAGAAAAACG
_	
SEQ2805	GATGAAAGCTATCTCTACCTTGCGATTAAAACAAAACCTGAAAAACTAAAAGAAAAACG
SEQ2806	Gatgaaagctatctctaccttgcgattaaaacaaaacctgaaaaactaaaagaaaacg
SEQ2807	GATGAAAGCTATCTCTACCTTGCGATTAAAACAAAACCTGAAAAACTAAAAGAAAAACG
SEQ2808	GATGAAAGCTATCTCTACCTTGCGATTAAAACAAAACCTGAAAAACTAAAAGAAAACG
- SEQ2809	GATGAAAGCTATCTCTACCTTGCGATTAAAACAAAACCTGAAAAACTAAAAGAAAAACG
SEQ2810	GATGAAAGCTATCTCTACCTTGCGATTAAAACAAAACCTGAAAAACTAAAAGAAAAACG
SE02811	
2EQ2811	GATGAAAGCTATCTCTACCTTGCGATTAAAACAAAACCTGAAAAACTAAAAGAAAAACG
•	
SEQ2801	
SEQ2802	
SEQ2803	TTATTACCAATAGATATTACACCAAAATCTGGTAGTAGAAAAATGAATG
SEQ2804	TTATTACCAATAGATATTACACCAAAATCTGGTAGTAGAAAAATGAATG
-	
SEQ2805	TTATTACCAATAGATATTACACCAAAATCTGGTAGTAGAAAAATGAATG
SEQ2806	TTATTACCAATAGATATTACACCAAAATCTGGTAGTAGAAAAATGAATG
SEQ2807	TTATTACCAATAGATATTACACCAAAATCTGGTAGTAGAAAAATGAATG
SEQ2808	TTATTACCAATAGATATTACACCAAAATCTGGTAGTAGAAAAATGAATG
SEQ2809	TTATTACCAATAGATATTACACCAAAATCTGGTAGTAGAAAAATGAATG
'SEQ2810	TTATTACCAATAGATATTACACCAAAATCTGGTAGTAGAAAAATGAATG
_	
SEQ2811	TTATTACCAATAGATATTACACCAAAATCTGGTAGTAGAAAAATGAATG
SEQ2801	
SEQ2802	
SEQ2803	ACATTTCTAAATCTAGTGACTTTGTATTGTCTATTGATCCAAATGGCAAGTCTGAATT
SEQ2804	ACATTTTCTAAATCTAGTGACTTTGTATTGTATTGATCCAAATGGCAAGTCTGAATT
SEQ2805	
	ACATTTTCTAAATCTAGTGACTTTGTATTGTCTATTGATCCAAATGGCAAGTCTGAATT
SEQ2806	ACATTTCTAAATCTAGTGACTTTGTATTGTCTATTGATCCAAATGGCAAGTCTGAATT
SEQ2807	ACATTTTCTAAATCTAGTGACTTTGTATTGTCTATTGATCCAAATGGCAAGTCTGAATT
SEQ2808	ACATTTCTAAATCTAGTGACTTTGTATTGTCTATTGATCCAAATGGCAAGTCTGAATT
SEQ2809	ACATTTCTAAATCTAGTGACTTTGTATTGTCTATTGATCCAAATGGCAAGTCTGAATT
SEQ2810	ACATTTCTAAATCTAGTGACTTTGTATTGTCTATTGATCCAAATGGCAAGTCTGAATT
SEQ2811	ACATTTCTAAATCTAGTGACTTTGTATTGTCTATTGATCCAAATGGCAAGTCTGAATT
022.022	ACATTITOTAMICIAGIGACTITGTATIGTCTATITGATCCAAATGGCAAGTCTGAATT
SEQ2801	
SEQ2802	
SEQ2803	TTTGTCCAAGAGCGCTATAATGCCTTAAAAGCGAACTATCTTCGACAGCTTAACGGTAA
SEQ2804	TTTGTCCAAGAGCGCTATAATGCCTTAAAAGCGAACTATCTTCGACAGCTTAACGGTAA
SEQ2805	TTTGTCCAAGAGCGCTATAATGCCTTAAAAGCGAACTATCTTCGACAGCTTAACGGTAA
SEQ2806	
***	TTTGTCCAAGAGCGCTATAATGCCTTAAAAGCGAACTATCTTCGACAGCTTAACGGTAA
SEQ2807	TTTGTCCAAGAGCGCTATAATGCCTTAAAAGCGAACTATCTTCGACAGCTTAACGGTAA
SEQ2808	TTTGTCCAAGAGCGCTATAACGCCTTAAAAGCGAACTATCTTCGACAGCTTAATGGTAA
SEQ2809	TTTGTCCAAGAGCGCTATAACGCCTTAAAAGCGAACTATCTTCGACAGCTTAATGGTAA
SEQ2810	TTTGTCCAAGAGCGCTATAATGCCTTAAAAGCGAACTATCTTCGACAGCTTAACGGTAA
SEQ2811	TTTGTCCAAGAGCGCTATAATGCCTTAAAAGCGAACTATCTTCGACAGCTTAACGGTAA
######################################	
SEQ2801	
SEQ2802	
SEQ2803	GATTTTTATGCTTTCCCACCAAAGAAGAACAGTAGTAATTTTGAGCAGATAAATATGGT
SEQ2804	GATTTTATGCTTTCCCACCAAAGAAGAACAGTAGTAATTTTGAGCAGATAAATATGGT
SEQ2805	GATTTTTATGCTTTCCCACCAAAGAAGAACAGTAGTAATTTTGAGCAGATAAATATGGT
SEQ2806	
	GATTTTTATGCTTTCCCACCAAAGAAGAACAGTAGTAATTTTGAGCAGATAAATATGGT
SEQ2807	GATTTTTATGCTTTCCCACCAAAGAAGAACAGTAGTAATTTTGAGCAGATAAATATGGT
SEQ2808	GATTTTTATGCTTTCCCACCAAAGAAGAACAGTAGTAATTTTGAGCAGATAAATATGGT
SEQ2809	GATTTTTATGCTTTCCCACCAAAGAAGAACAGTAGTAATTTTGAGCAGATAAATATGGT
SEQ2810	GATTTTTATGCTTTCCCACCAAAGAAGAACAGTAGTAATTTTGAGCAGATAAATATGGT
SEQ2811	GATTTTTATGCTTTCCCACCAAAGAAGAACAGTAGTAATTTTGAGCAGATAAATATGGT
~~ X~~~	CALLE TATOCITICO CACCAMAGAAGAACAGTAGTATTTTGAGCAGATAAATATGGT
CT-00055	
SEQ2801	
SEQ2802	
SEQ2803	TTGAGAAATACAAAGATTGTTGAAGACATGGAAAAAGTAAAAGCAACAGAGAGGTTCTT
SEQ2804	TTGAGAAATACAAAGATTGTTGAAGACATGGAAAAAGTAAAAGCAACAGAGAGGTTCTT
-	The state of the s



## Table 26: C mparative Sequences relating to SAG1552 (conserved hypothetical pr tein)







SEQ2801	
SEQ2802	
SEQ2803	AAACCTTTTTAAAAGACTCCTATTATGTATTAAGAAAGAA
SEQ2804	AAACCTTTTTAAAAGACTCCTATTATAGTATTAAGAAAGA
SEQ2805	AAACCTTTTTAAAAGA
SEQ2806	AAACCTTTTTAAAAGACTCCTATTATGTATTAAGAAAGA
SEQ2807	AAACCTTTTTAAAAGACT
SEQ2808	AAACCTTTTTAAAAGACTCCTATTATAGT
SEQ2809	AAACCTTTTTAAAAGACTCCTATTATAGTATTAAGAAAG
SEQ2810	AAACCTTTTTAAAAGACTCCTATTATAGTATTAAG
SEQ2811	AAACCTTTTTAAAAGACTCCTATTATAGTATTAAGAAAGA
•	
SEQ2801	
SEQ2802	
SEQ2803	
SEQ2804	GAACATATGGTCCA
SEQ2805	
SEQ2806	
SEQ2807	*============
SEQ2808	=======================================
SEQ2809	
SEQ2810	
SEQ2811	= = = = = = = = = = = = = = = = = = =

#### >SEQ ID NO 2850:62 1169NT frame: 1

FVVKGDTVLHKPTNKPFVVKGVDVESSLAGYHHNDFPITQKTYREWFHLISNMGANTVRV KVPMNVAFYDALYHHNKASKRPLYLLQGIRIDSYRNNASITAFNDNYRGYLKREAKGVVD ILHGRKQVWNTDFGSRHYHYDLSPWVLGYVVGDDWNSGTVAYTNHQEKKTQYKGRYFKTS AAANPFEVMLAQVMDELTHYETAKYGWQHLISFSNSPTTDPFRYRKPFEAQAPKYVQLNV ENIQANSNVKAGIFAAYKAIDFHPRYKDYLLFDKENISKEDRQKIKELSLSQGYVKLLNA YHKIPVLVTGYGYSTARGIAQKEIDKRPLPINEKEQGQRLLEDYESFISSGSFGATINAW QDDWNARAWNTSFATNKHSQFLWGDAQVFNQGYGLLGFKNAKHHYQVDGKRGKGEWKHPL MTSATGDDLYASSDESYLYLAIKTKPEKLKEKRLLPIDITPKSGSRKMNGSKVTFSKSSD FVVLSIDPNGKSELFVQERYNALKANYLRQLNGKDFYAFPPKKNSSNFEQINMVLRNTKIV EDMEKVKATERFLPTHPTGLLKTGTIDRHQKTFDSQTDISFGKDFIEVRIPWQLLNFSDP SSQKIHDDYFKHYGVKELEIESIALGLGANSKENTLIKMADYRLKNWERPDTKTFLKDSY YSI.ER

### >SEQ ID NO 2851:62\_18RS21 frame: 1

KGLLKENTRTNFVVKGDTVLHKPTNKPFVVKGVDVESSLAGYHHNDFPITQKTYREWFHL ISNMGANTVRVKVPMNVAFYDALYHHNKASKRPLYLLQGIRIDSYRNNASITAFNDNYRG YLKREAKGVVDILHGRKQVWNTDLGSRHYHYDLSPWVLGYVVGDDWNSGTVAYTNHQEKK TQYKGRYFKTSVAANPFEVMLAQVMDELTHYETAKYGWQHLISFSNSPTTDPFHYRKPFE AQAPKYVQLNVENIQANSNVKAGMFAAYKAIDFHPRYKDYLLFDKENISKEDRQKIKELS LSQGYVKLLNAYHKIPVLVTGYGYSTARGIAQKEIDKRPLPINEKEQGQRLLEDYESFIS SGSFGATINAWQDDWNARAWNTSFATNKHSQFLWGDAQVFNQGYGLLGFKNAKHHYQVDG KRGKGEWKHPLMTSATGDDLYASSDESYLYLAIKTKPEKLKEKRLLPIDITPKSGSRKMN GSKVTFSKSSDFVLSIDPNGKSELFVQERYNALKANYLRQLNGKDFYAFPPKKNSSNFEQ INMVLRNTKIVEDMEKVKATERFLPTHPTGLLKTGTTDRHQKTFDSQTDISFGKDFIEVR IPWQLLNFSDPSSQKIHDDYFKHYGVKELEIESIALGLGANSKENTLIKMADYRLKNWER PDTKTFLKDSYYVLRK

### >SEQ ID NO 2852:62\_2603 frame: 3

LKENTRTNFVVKGDTVLHKPTNKPFVVKGVDVESSLAGYHHNDFPITQKTYREWFHLISN MGANTVRVKVPMNVAFYDALYHHNKASKRPLYLLQGIRIDSYRNNASITAFNDNYRGYLK REAKGVVDILHGRKQVWNTDLGSRHYHYDLSPWVLGYVVGDDWNSGTVAYTNHQEKKTQY KGRYFKTSVAANPFEVMLAQVMDELTHYETAKYGWQHLISFSNSPTTDPFHYRKPFEAQA PKYVQLNVENIQANSNVKAGMFAAYKAIDFHPRYKDYLLFDKENISKEDRQKIKELSLSQ GYVKLLNAYHKIPVLVTGYGYSTARGIAQKEIDKRPLPINEKEQGQRLLEDYESFISSGS FGATINAWQDDWNARAWNTSFATNKHSQFLWGDAQVFNQGYGLLGFKNAKHYQVDGKRG KGEWKHPLMTSATGDDLYASSDESYLYLAIKTKPEKLKEKRLLPIDITPKSGSRKMNGSK VTFSKSSDFVLSIDPNGKSELFVQERYNALKANYLRQLNGKDFYAFPPKKNSSNFEQINM VLRNTKIVEDMEKVKATERFLPTHPTGLLKTGTTDRHQKTFDSQTDISFGKDFIEVRIPW QLLNFSDPSSQKIHDDYFKHYGVKELEIESIALGLGANSKENTLIKMADYRLKNWERPDT KTFLKDSYYSIKKEWSKERERTYGP



# Table 26: C mparative Sequences relating to SAG1552 (conserved hypothetical pr tein)

### >SEQ ID NO 2853:62\_A909 frame: 1

KGLLKENTRTNFVVKGDTVLHKPTNKPFVVKGVDVESSLAGYHHNDFPITQKTYREWFHL ISNMGANTVRVKVPMNVAFYDALYHHNKASKRPLYLLQGIRIDSYRNNASITAFNDNYRG YLKREAKGVVDILHGRKQVWNTDLGSRHYHYDLSPWVLGYVVGDDWNSGTVAYTNHQEKK TQYKGRYFKTSVAANPFEVMLAQVMDELTHYETAKYGWQHLISFSNSPTTDPFHYRKPFE AQAPKYVQLNVENIQANSNVKAGMFAAYKAIDFHPRYKDYLLFDKENISKEDRQKIKELS LSQGYVKLINAYHKIPVLVTGYGYSTARGIAQKEIDKRPLPINEKEQGQRLLEDYESFIS SGSFGATINAWQDDWNARAWNTSFATNKHSQFLWGDAQVFNQGYGLLGFKNAKHHYQVDG KRGKGEWKHPLMTSATGDDLYASSDESYLYLAIKTKPEKLKEKRLLPIDITPKSGSRKMN GSKVTFSKSDFVLSIDPNGKSELFVQERYNALKANYLRQLNGKDFYAFPPKKNSSNFEQ INMVLRNTKIVEDMEKVKATERFLPTHPTGLLKTGTTDRHQKTFDSQTDISFGKDFIEVR IPWQLLNFSDPSSQRIHDDYFKHYGVKELEN.EPLL.D.VLIAKKTH.RWQIIV.KIGR DPIPKFF.K

### >SEQ ID NO 2854:62 A909 frame: 1

KGLLKENTRTNFVVKGDTVLHKPTNKPFVVKGVDVESSLAGYHHNDFPITQKTYREWFHL ISNMGANTVRVKVPMNVAFYDALYHHNKASKRPLYLLQGIRIDSYRNNASITAFNDNYRG YLKREAKGVVDILHGRKQVWNTDLGSRHYHYDLSPWVLGYVVGDDWNSGTVAYTNHQEKK TQYKGRYFKTSVAANPFEVMLAQVMDELTHYETAKYGWQHLISFSNSPTTDPFHYRKPFE AQAPKYVQLNVENIQANSNVKAGMFAAYKAIDFHPRYKDYLLFDKENISKEDRQKIKELS LSQGYVKLLNAYHKIPVLVTGYGYSTARGIAQKEIDKRPLPINEKEQGQRLLEDYESFIS SGSFGATINAWQDDWNARAWNTSFATTNKHSQFLWGDAQVFNQGYGLLGFKNAKHHYQVDG KRGKGEWKHPLMTSATGDDLYASSDESYLYLAIKTKPEKLKEKRLLPIDITPKSGSRKMN GSKVTFSKSSDFVLSIDPNGKSELFVQERYNAKANYLRQLNGKDFYAFPPKKNSSNFEQ INMVLRNTKIVEDMEKVKATERFLPTHPTGLLKTGTTDRHQKTFDSQTDISFGKDFIEVR IPWQLLNFSDPSSQRIHDDYFKHYGVKELEN.EPLL.D.VLIAKKTH.RWQIIV.KIGR DPIFKPF.K

#### >SEQ ID NO 2855:62\_CJB110 frame: 1

YYFDGSLYLPKGLLKENTRTNFVVKGDTVLHKPTNKPFVVKGVDVESSLAGYHHNDFPIT QKTYREWFHLISNMGANTVRVKVPMNVAFYDALYHHNKASKRPLYLLQGIRIDSYRNNAS ITAFNDNYRGYLKREAKGVVDILHGRKQVWNTDFGSRHYHYDLSPWVLGYVVGDDWNSGT VAYTNHQEKKTQYKGRYFKTSVAANPFEVMLAQVMDELTHYETAKYGWQHLISFSNSPTT DPFHYRKPFEAQAPKYVQLNVENIQANSNVKAGMFAAYKAIDFHPRYKDYLLFDKENISK EDRQKIKELSLSQGYVKLLNAYHKIPVLVTGYGYSTARGIAQKEIDKRPLPINEKEQGQR LLEDYESFISSGSFGATINAWQDDWNARAWNTSFATNKHNQFLWGDAQVFNQGYGLLGFK NAKHHYQVDGKRGKGEWKHPLMTSATGDDLYASDESYLYLAIKTKPEKLKEKRLLPIDI TPKSGSRKMNGSKVTFSKSSDFVLSIDPNGKSELFVQERYNALKANYLRQLNGKDFYAFP PKKNSSNFEQINMVLRNTKIVEDMEKVKATERFLPTHPTGLLKTGTTDRHQKTFDSQTDI SFGKDFIEVRIPWQLLNFSDPSSQKIHDDYFKHYGVKELEIESIALGLGANSKENTLIKM ADYRLKNWERPDTKTFLKDSYYVLRK

### >SEQ ID NO 2856:62\_COH1 frame: 2

LPQGLLKENTRTNFVVKGDTVLHKPTNKPFVVKGVDVESSLAGYHHNDFPITQKTYREWF HLISNMGANTVRVKVPMNVAFYDALYHHNKESKRPLYLLQGIRIDSYRNNASITAFNDNY RGYLKREAKGVVDILHGRKQVWNTDFGSRHYHYDLSPWVLGYVVGDDWNSGTVAYTNHQE KKTQYKGRYFKTSVAANPFEVMLAQVMDELTHYETAKYGWQHLISFSNSPTTDPFHYRKP FEAQAPKYVQLNVENIQANSNVKAGMFAAYKAIDFHPRYKDYLLFDKENISKEDRQKIKE LSLSQGYVKLLNAYHKIPVLVTGYGYSTARGIAQKEIDKRPLPINEKEQGQRLLEDYESF ISSGSFGATINAWQDDWNARAWNTSFATNKHSQFLWGDAQVFNQGYGLLGFKNAKHHYQV DGKRGKGEWKHPLMTSATGDDLYASSDESYLYLAIKTKPEKLKEKRLLPIDITPKSGSRK MNGSKVTFSKSSDFVLSIDPNGKSELFVQERYNALKANYLRQLNGKDFYAFPPKKNSSNF EQINMVLRNTKIVEDMEKVKATERFLPTHPTGLLKTGTTDRHQKTFDSQPDISFGKDFIE VRIPWQLLNFSDPSSQKIHDDYFKHYGVKELEIESIALGLGANSKENTLIKMADYRLKNW ERPDTKTFLKD

#### >SEQ ID NO 2857:62 H36B frame: 2

RGLLKENTRTNFVVKGDTVLHKPTNKPFVVKGVDVESSLAGYHHNDFPITQKTYREWFHL ISNMGANTVRVKVPMNVAFYDALYHHNKASKRPLYLLQGIRIDSYRNNASITAFNDNYRG YLKREAKGVVDILHGRKQVWNTDFGSSHYHYDLSPWVLGYVVGDDGHSGTVALY

### >SEQ ID NO 2858:62\_JM9130013 frame: 3

FVVKGDTVLHKPTNKPFVVKGVDVESSLAGYHHNDFPITQKTYREWFHLISNMGANTVRV KVPMNVAFYDALYHHNKASKRPLYLLQGIRIDSYRNNASITAFNDNYRGYLKREAKGVVD ILHGRKQVWNTDFGSSHYHYDLSPWVLGYVVGDDWNSGTVAYTNHQEKKTQYKGRYFKTS VAANPFEVMLAQVMDELTHYETAKYGWQHLISFSNSPTTDPFHYRKPFEAQAPKYVQLNV ENIQANSNVKAGMFAAYKAIDFHPRYKDYLLFDKENISKEDRQKIKELSLSQGYVKLLNA YHKIPVLVTGYGYSTARGIAQKEIDKRPLPINEKEQGQRLLEDYESFISSGSFGATINAW



QDDWNARVWNTSFATNKHSQFLWGDAQVFNQGYGLLGFKNAKHHYQVDGKRGKEEWKHPL MTSATGDDLYASSDESYLYLAIKTKPEKLKEKRLLPIDITPKSGSRKMNGSKVTFSKSSD FVLSIDPNGKSELFVQERYNALKANYLRQLNGKDFYAFPPKKNSSNFEQINMVLRNTKIV EDMEKVKATERFLPTHPTGLLKTGTTDRHQKTFDSQTDISFGKDFIEVRIPWQLLNFSDP SSQKIHDDYFKHYGVKELEIESIALGLGANSKENTLIKMADYRLKNWERPDTKTFLKDSY YSIKK

#### >SEQ ID NO 2859:62\_M732 frame: 2

TRTNFVVKGDTVLHKPTNKPFVVKGVDVESSLAGYHHNDFPITQKTYREWFHLISNMGAN TVRVKVPMNVAFYDALYHHNKESKRPLYLLQGIRIDSYRNNASITAFNDNYRGYLKREAK GVVDILHGRKQVWNTDFGSRHYHYDLSPWVLGYVVGDDCNSGTVAYTNHQEKKTQYKGRY FKTSVAANPFEVMLAQVMDELTHYETAKYGWQHLISFSNSPTTDPFHYRKPFEAQAPKYV QLNVENIQANSNVKAGMFAAYKAIDFHPRYKDYLLFDKENISKEDRQKIKELSLSQGYVK LLNAYHKIPVLVTGYGYSTARGIAQKEIDKRPLPINEKEQGQRLLEDYESFISSGSFGAT INAWQDDWNARAWNTSFATNKHSQFLWGDAQVFNQGYGLLGFKNAKHHYQVDGKRGKGEW KHPLMTSATGDDLYASSDESYLYLAIKTKPEKLKEKRLLPIDITPKSGSRKMNGSKVTFS KSDFVLSIDPNGKSELFVQERYNALKANYLRQLNGKDFYAFPPKKNSSNFEQINMVLRN TKIVEDMEKVKATERFLPTHPTGLLKTGTTDRHQKTFDSQTDISFGKDFIEVRIPWQLLN FSDPSSQKIHDDYFKHYGVKELEIESIALGLGANSKENTLIKMADYRLKNWERPDTKTFL KDSYYSIK

#### >SEQ ID NO 2860:62 M781 frame: 1

FDGSLYLPQGLLKENTRTNFVVKGDTVLHKPTNKPFVVKGVDVESSLAGYHHNDFPITQK
TYREWFHLISNMGANTVRVKVPMNVAFYDALYHHNKESKRPLYLLQGIRIDSYRNNASIT
AFNDNYRGYLKREAKGVVDILHGRKQVWNTDFGSRHYHYDLSPWVLGYVVGDDWNSGTVA
YTNHQEKKTQYKGRYFKTSVAANPFEVMLAQVMDELTHYETAKYGWQHLISFSNSPTTDP
FHYRKPFEAQAPKYVQLNVENIQANSNVKAGMFAAYKAIDFHPRYKDYLLFDKENISKED
RQKIKELSLSQGYVKLLNAYHKIPVLVTGYGYSTARGIAQKEIDKRPLPINEKEQGQRLL
EDYESFISSGSFGATINAWQDDWNARAWNTSFATNKHSQFLWGDAQVFNQGYGLLGFKNA
KHHYQVDGKRGKGEWKHPLMTSATGDDLYASSDESYLYLAIKTKPEKLKEKRLLPIDITP
KSGSRKMNGSKVTFSKSSDFVLSIDPNGKSELFVQERYNALKANYLRQLNGKDFYAFPPK
KNSSNFEQINMVLRNTKIVEDMEKVKATERFLPTHPTGLLKTGTTDRHQKTFDSQTDISF
GKDFIEVRIPWQLLNFSDPSSQKIHDDYFKHYGVKELEIESIALGLGANSKENTLIKMAD
YRLKNWERPDTKTFLKDSYYSIKKEW

SEQ2850	FVVKGDTVLHKPTNKPFVVKGVDVESSLAGYHHNDFPIT
SEQ2851	KGLLKENTRTNFVVKGDTVLHKPTNKPFVVKGVDVESSLAGYHHNDFPIT
SEQ2852	LKENTRTNFVVKGDTVLHKPTNKPFVVKGVDVESSLAGYHHNDFPIT
SEQ2853	KGLLKENTRTNEVVKGDTVLHKPTNKPEVVKGVDVESSLAGYHHNDFPIT
SEQ2854	KGLLKENTRTNFVVKGDTVLHKPTNKPFVVKGVDVESSLAGYHHNDFPIT
SEQ2855	YFDGSLYLPKGLLKENTRTNFVVKGDTVLHKPTNKPFVVKGVDVESSLAGYHHNDFPIT
SEQ2856	LPQGLLKENTRTNFVVKGDTVLHKPTNKPFVVKGVDVESSLAGYHHNDFPTT
SEQ2857	RGLLKENTRTNFVVKGDTVLHKPTNKPFVVKGVDVESSLAGYHHNDFPTT
SEQ2858	FVVKGDTVLHKPTNKPFVVKGVDVESSLAGYHHNDFPIT
SEQ2859	TRTNFVVKGDTVLHKPTNKPFVVKGVDVESSLAGYHHNDFPIT
SEQ2860	-FDGSLYLPQGLLKENTRTNFVVKGDTVLHKPTNKPFVVKGVDVESSLAGYHHNDFPIT
SEQ2850	QKTYREWFHLISNMGANTVRVKVPMNVAFYDALYHHNKASKRPLYLLQGIRIDSYRNNAS
SEQ2851	QKTYREWFHLISNMGANTVRVKVPMNVAFYDALYHHNKASKRPLYLLQGIRIDSYRNNAS
SEQ2852	QKTYREWFHLISNMGANTVRVKVPMNVAFYDALYHHNKASKRPLYLLQGIRIDSYRNNAS
SEQ2853	QKTYREWFHLISNMGANTVRVKVPMNVAFYDALYHHNKASKRPLYLLQGIRIDSYRNNAS
SEQ2854	QKTYREWFHLISNMGANTVRVKVPMNVAFYDALYHHNKASKRPLYLLQGIRIDSYRNNAS
SEQ2855	QKTYREWFHLISNMGANTVRVKVPMNVAFYDALYHHNKASKRPLYLLQGIRIDSYRNNAS
SEQ2B56	QKTYREWFHLISNMGANTVRVKVPMNVAFYDALYHHNKESKRPLYLLQGIRIDSYRNNAS
SEQ2857	QKTYREWFHLISNMGANTVRVKVPMNVAFYDALYHHNKASKRPLYLLQGIRIDSYRNNAS
SEQ2858	QKTYREWFHLISNMGANTVRVKVPMNVAFYDALYHHNKASKRPLYLLQGIRIDSYRNNAS
SEQ2859	QKTYREWFHLISNMGANTVRVKVPMNVAFYDALYHHNKESKRPLYLLQGIRIDSYRNNAS
SEQ2860	QKTYREWFHLISNMGANTVRVKVPMNVAFYDALYHHNKESKRPLYLLQGIRIDSYRNNAS
SEQ2850	ITAFNDNYRGYLKREAKGVVDILHGRKQVWNTDFGSRHYHYDLSPWVLGYVVGDDWNSGT
SEQ2851	ITAFNDNYRGYLKREAKGVVDILHGRKQVWNTDLGSRHYHYDLSPWVLGYVVGDDWNSGT
SEQ2852	ITAFNDNYRGYLKREAKGVVDILHGRKQVWNTDLGSRHYHYDLSPWVLGYVVGDDWNSGT
SEQ2853	ITAFNDNYRGYLKREAKGVVDILHGRKQVWNTDLGSRHYHYDLSPWVLGYVVGDDWNSGT
SEQ2854	ITAFNDNYRGYLKREAKGVVDILHGRKQVWNTDLGSRHYHYDLSPWVLGYVVGDDWNSGT
SEQ2855	ITAFNDNYRGYLKREAKGVVDILHGRKQVWNTDFGSRHYHYDLSPWVLGYVVGDDWNSGT
SEQ2856	ITAFNDNYRGYLKREAKGVVDILHGRKQVWNTDFGSRHYHYDLSPWVLGYVVGDDWNSGT
SEQ2857	ITAFNDNYRGYLKREAKGVVDILHGRKQVWNTDFGSSHYHYDLSPWVLGYVVGDDGHSGT
SEQ2858	ITAFNDNYRGYLKREAKGVVDILHGRKQVWNTDFGSSHYHYDLSPWVLGYVVGDDWNSGT



# Table 28: Comparative Sequences relating to SAG1552 (conserved hypothetical pr tein)

SEQ2859 SEQ2860 SEQ2850 SEQ2851 SEQ2852 SEQ2853 SEQ2854 SEQ2855 SEQ2856 SEQ2856	ITAFNDNYRGYLKREAKGVVDILHGRKQVWNTDFGSRHYHYDLSPWVLGYVVGDDCNSGT ITAFNDNYRGYLKREAKGVVDILHGRKQVWNTDFGSRHYHYDLSPWVLGYVVGDDWNSGT VAYTNHQEKKTQYKGRYFKTSAAANPFEVMLAQVMDELTHYETAKYGWQHLISFSNSPTT VAYTNHQEKKTQYKGRYFKTSVAANPFEVMLAQVMDELTHYETAKYGWQHLISFSNSPTT VAYTNHQEKKTQYKGRYFKTSVAANPFEVMLAQVMDELTHYETAKYGWQHLISFSNSPTT VAYTNHQEKKTQYKGRYFKTSVAANPFEVMLAQVMDELTHYETAKYGWQHLISFSNSPTT VAYTNHQEKKTQYKGRYFKTSVAANPFEVMLAQVMDELTHYETAKYGWQHLISFSNSPTT VAYTNHQEKKTQYKGRYFKTSVAANPFEVMLAQVMDELTHYETAKYGWQHLISFSNSPTT VAYTNHQEKKTQYKGRYFKTSVAANPFEVMLAQVMDELTHYETAKYGWQHLISFSNSPTT VAYTNHQEKKTQYKGRYFKTSVAANPFEVMLAQVMDELTHYETAKYGWQHLISFSNSPTT VAYTNHQEKKTQYKGRYFKTSVAANPFEVMLAQVMDELTHYETAKYGWQHLISFSNSPTT
SEQ2858	VAYTNHQEKKTQYKGRYFKTSVAANPFEVMLAQVMDELTHYETAKYGWQHLISFSNSPTT
SEQ2859	VAYTNHQEKKTQYKGRYFKTSVAANPFEVMLAQVMDELTHYETAKYGWQHLISFSNSPTT
SEQ2860	VAYTNHQEKKTQYKGRYFKTSVAANPFEVMLAQVMDELTHYETAKYGWQHLISFSNSPTT
J-2-000	
SEQ2850	PFRYRKPFEAQAPKYVQLNVENIQANSNVKAGIFAAYKAIDFHPRYKDYLLFDKENISK
SEQ2851	PFHYRKPFEAQAPKYVQLNVENIQANSNVKAGMFAAYKAIDFHPRYKDYLLFDKENISK
SEQ2852	PFHYRKPFEAQAPKYVQLNVENIQANSNVKAGMFAAYKAIDFHPRYKDYLLFDKENISK
SEQ2853	PFHYRKPFEAQAPKYVQLNVENIQANSNVKAGMFAAYKAIDFHPRYKDYLLFDKENISK
SEQ2854	PFHYRKPFEAQAPKYVQLNVENIQANSNVKAGMFAAYKAIDFHPRYKDYLLFDKENISK
SEQ2855	PFHYRKPFEAQAPKYVQLNVENIQANSNVKAGMFAAYKAIDFHPRYKDYLLFDKENISK
_	
SEQ2856	PFHYRKPFEAQAPKYVQLNVENIQANSNVKAGMFAAYKAIDFHPRYKDYLLFDKENISK
SEQ2857	DEUD PO DE A A A DEVITAT MUDIT A AMONTO A CHER A VER A DEUD PEROLET E PRESENT OF
SEQ2858	PFHYRKPFEAQAPKYVQLNVENIQANSNVKAGMFAAYKAIDFHPRYKDYLLFDKENISK
SEQ2859	PFHYRKPFEAQAPKYVQLNVENIQANSNVKAGMFAAYKAIDFHPRYKDYLLFDKENISK
SEQ2860	PFHYRKPFEAQAPKYVQLNVENIQANSNVKAGMFAAYKAIDFHPRYKDYLLFDKENISK
SEQ2850	DRQKIKELSLSQGYVKLLNAYHKIPVLVTGYGYSTARGIAQKEIDKRPLPINEKEQGQR
SEQ2851	DRQKIKELSLSQGYVKLLNAYHKIPVLVTGYGYSTARGIAQKEIDKRPLPINEKEQGQR
SEQ2852	DRQKIKELSLSQGYVKLLNAYHKIPVLVTGYGYSTARGIAQKEIDKRPLPINEKEQGQR
SEQ2853	DRQKIKELSLSQGYVKLLNAYHKIPVLVTGYGYSTARGIAQKEIDKRPLPINEKEQGQR
SEQ2854	DRQKIKELSLSQGYVKLLNAYHKIPVLVTGYGYSTARGIAQKEIDKRPLPINEKEQGQR
SEQ2855	DRQKIKELSLSQGYVKLLNAYHKIPVLVTGYGYSTARGIAQKEIDKRPLPINEKEQGQR
SEQ2856	DRQKIKELSLSQGYVKLLNAYHKIPVLVTGYGYSTARGIAQKEIDKRPLPINEKEQGQR
SEQ2857	
SEQ2858	DRQKIKELSLSQGYVKLLNAYHKIPVLVTGYGYSTARGIAQKEIDKRPLPINEKEQGQR
SEQ2859	drqkikelslsqgyvkllnayhkipvlvtgygystargiaqkeidkrplpinekeqgqr
SEQ2860	DRQKIKELSLSQGYVKLLNAYHKIPVLVTGYGYSTARGIAQKEIDKRPLPINEKEQGQR
SEQ2850	LEDYESFISSGSFGATINAWQDDWNARAWNTSFATNKHSQFLWGDAQVFNQGYGLLGFK
'SEQ2851	LEDYESFISSGSFGATINAWQDDWNARAWNTSFATNKHSQFLWGDAQVFNQGYGLLGFK
SEQ2852	LEDYESFISSGSFGATINAWQDDWNARAWNTSFATNKHSQFLWGDAQVFNQGYGLLGFK
SEQ2853	LEDYESFISSGSFGATINAWQDDWNARAWNTSFATNKHSQFLWGDAQVFNQGYGLLGFK
SEQ2854	LEDYESFISSGSFGATINAWQDDWNARAWNTSFATNKHSQFLWGDAQVFNQGYGLLGFK
SEQ2855	LEDYESFISSGSFGATINAWQDDWNARAWNTSFATNKHNQFLWGDAQVFNQGYGLLGFK
SEQ2856	LEDYESFISSGSFGATINAWQDDWNARAWNTSFATNKHSQFLWGDAQVFNQGYGLLGFK
SEQ2857	
SRQ2858	LEDYESFISSGSFGATINAWQDDWNARVWNTSFATNKHSQFLWGDAQVFNQGYGLLGFK
SEQ2859	LEDYESFISSGSFGATINAWQDDWNARAWNTSFATNKHSQFLWGDAQVFNQGYGLLGFK
\$EQ2860	LEDYESFISSGSFGATINAWQDDWNARAWNTSFATNKHSQFLWGDAQVFNQGYGLLGFK
SEQ2850	AKHHYQVDGKRGKGEWKHPLMTSATGDDLYASSDESYLYLAIKTKPEKLKEKRLLPIDI
SEQ2851	AKHHYOVDGKRGKGEWKHPLMTSATGDDLYASSDESYLYLAIKTKPEKLKEKRLLPIDI
SEQ2852	AKHHYQVDGKRGKGEWKHPLMTSATGDDLYASSDESYLYLAIKTKPEKLKEKRLLPIDI
SEQ2853	AKHHYQVDGKRGKGEWKHPLMTSATGDDLYASSDESYLYLAIKTKPEKLKEKRLLPIDI
SEQ2854	AKHHYQVDGKRGKGEWKHPLMTSATGDDLYASSDESYLYLAIKTKPEKLKEKRLLPIDI
SEQ2855	AKHHYQVDGKRGKGEWKHPLMTSATGDDLYASSDESYLYLAIKTKPEKLKEKRLLPIDI
SEQ2856	AKHHYQVDGKRGKGEWKHPLMTSATGDDLYASSDESYLYLAIKTKPEKLKEKRLLPIDI
SEQ2857	
SEQ2858	AKHHYQVDGKRGKEEWKHPLMTSATGDDLYASSDESYLYLAIKTKPEKLKEKRLLPIDI
SEQ2859	AKHHYQVDGKRGKGEWKHPLMTSATGDDLYASSDESYLYLAIKTKPEKLKEKRLLPIDI  AKHHYQVDGKRGKGEWKHPLMTSATGDDLYASSDESYLYLAIKTKPEKLKEKRLLPIDI
SEQ2860	AKHHYQVDGKRGKGEWKHPLMTSATGDDLYASSDESYLYLAIKTKPEKLKEKRLLPIDI
SEQ2850	PKSGSRKMNGSKVTFSKSSDFVLSIDPNGKSELFVQERYNALKANYLROLNGKDFYAFP
SEQ2851	PKSGSRKMNGSKVTFSKSSDFVLSIDPNGKSELFVQERYNALKANYLROLNGKDFYAFP
SEQ2852	PKSGSRKMNGSKVTFSKSSDFVLSIDPNGKSELFVQERYNALKANYLRQLNGKDFYAFP
SEQ2853	PKSGSRKMNGSKVTFSKSSDFVLSIDPNGKSELFVQERYNALKANYLRQLNGKDFYAFP
SEQ2854	PKSGSRKMNGSKVTFSKSSDFVLSIDPNGKSELFVQERYNALKANYLROLNGKDFYAFP
SEQ2855	PKSGSRKMNGSKVTFSKSSDFVLSIDPNGKSELFVQERYNALKANYLRQLNGKDFYAFP
2485444	





SEQ2856	PKSGSRKMNGSKVTFSKSSDFVLSIDPNGKSELFVQERYNALKANYLRQLNGKDFYAFP
SEQ2857	
SEQ2858	PKSGSRKMNGSKVTFSKSSDFVLSIDPNGKSELFVQERYNALKANYLRQLNGKDFYAFP
SEQ2859	PKSGSRKMNGSKVTFSKSSDFVLSIDPNGKSELFVQERYNALKANYLRQLNGKDFYAFP
SEQ2860	PKSGSRKMNGSKVTFSKSSDFVLSIDPNGKSELFVQERYNALKANYLRQLNGKDFYAFP
SEQ2850	KKNSSNFEQINMVLRNTKIVEDMEKVKATERFLPTHPTGLLKTGTIDRHQKTFDSQTDI
SEQ2851	KKNSSNFEQINMVLRNTKIVEDMEKVKATERFLPTHPTGLLKTGTTDRHQKTFDSQTDI
SEQ2852	KKNSSNFEQINMVLRNTKIVEDMEKVKATERFLPTHPTGLLKTGTTDRHQKTFDSQTDI
SEQ2853	KKNSSNFEQINMVLRNTKIVEDMEKVKATERFLPTHPTGLLKTGTTDRHQKTFDSQTDI
SEQ2854	KKNSSNFEQINMVLRNTKIVEDMEKVKATERFLPTHPTGLLKTGTTDRHQKTFDSQTDI
SEQ2855	KKNSSNFEQINMVLRNTKIVEDMEKVKATERFLPTHPTGLLKTGTTDRHQKTFDSQTDI
SEQ2856	KKNSSNFEQINMVLRNTKIVEDMEKVKATERFLPTHPTGLLKTGTTDRHQKTFDSQPDI
SEQ2857	
SEQ2858	KKNSSNFEQINMVLRNTKIVEDMEKVKATERFLPTHPTGLLKTGTTDRHQKTFDSQTDI
SEQ2859	KKNSSNFEQINMVLRNTKIVEDMEKVKATERFLPTHPTGLLKTGTTDRHQKTFDSQTDI
SEQ2860	KKNSSNFEQINMVLRNTKIVEDMEKVKATERFLPTHPTGLLKTGTTDRHQKTFDSQTDI
SEQ2850	FGKDFIEVRIPWQLLNFSDPSSQKIHDDYFKHYGVKELEIESIALGLGANSKENTLIKM
SEQ2851	FGKDFIEVRIPWQLLNFSDPSSQKIHDDYFKHYGVKELEIESIALGLGANSKENTLIKM
SEQ2852	FGKDFIEVRIPWQLLNFSDPSSQKIHDDYFKHYGVKELEIESIALGLGANSKENTLIKM
SEQ2853	FGKDFIEVRIPWQLLNFSDPSSQRIHDDYFKHYGVKELENEPLLDVLIAKKTHRWQIIV
SEQ2854	FGKDFIEVRIPWQLLNFSDPSSQRIHDDYFKHYGVKELENEPLLDVLIAKKTHRWQIIV
SEQ2855	FGKDFIEVRIPWQLLNFSDPSSQKIHDDYFKHYGVKELEIESIALGLGANSKENTLIKM
SEQ2856	FGKDFIEVRIPWQLLNFSDPSSQKIHDDYFKHYGVKELEIESIALGLGANSKENTLIKM
SEQ2857	
SEQ2858	FGKDFIEVRIPWQLLNFSDPSSQKIHDDYFKHYGVKELEIESIALGLGANSKENTLIKM
SEQ2859	FGKDFIEVRIPWQLLNFSDPSSQKIHDDYFKHYGVKELEIESIALGLGANSKENTLIKM
SEQ2860	FGKDFIEVRIPWQLLNFSDPSSQKIHDDYFKHYGVKELEIESIALGLGANSKENTLIKM
SEQ2850	DYRLKNWERPDTKTFLKDSYYSIER
SEQ2851	DYRLKNWERPDTKTFLKDSYYVLRK
SEQ2852	DYRLKNWERPDTKTFLKDSYYSIKKEWSKERERTYGP
SEQ2853	IGRDPIPKPFK
SEQ2854	IGRDPIPKPFK
SEQ2855	DYRLKNWERPDTKTFLKDSYYVLRK
SEQ2856	DYRLKNWERPDTKTFLKD
SEQ2857	
SEQ2858	DYRLKNWERPDTKTFLKDSYYSIKK
SEQ2859	DYRLKNWERPDTKTFLKDSYYSIK
SEQ2860	DYRLKNWERPDTKTFLKDSYYSIKKEW



### Table 29: C mparative Sequences relating to SAG1641 (YaeC family pr tein)

### SEQ ID NO. 2902: SAG1641 FROM THE 1169NT1 GBS TYPE V STRAIN (REVERSE COMPLEMENT)

### Table 29: Comparative Sequences relating t SAG1641 (YaeC family protein)

### SEQ ID NO. 2907: SAG1641 FROM THE COH1 GBS TYPE III STRAIN (REVERSE COMPLEMENT)

### SEQ ID NO. 2908: SAG1641 FROM THE H36b GBS TYPE Ib STRAIN

AAGAAGTTTCAGCAAGCTCAACTTCAAGTAAAGTTGTTAAAGTTGGTGTTATGACCTTTTCTGACACTGAAAAAGCAC
GTTGGGATAAAATTGAAAAGCTAGTAGGTGATAAAGCTAAAATCAAATTTACAGAATTTACAGATTATACACAACCAA
ATCAAGCGACCCAATAAGGATGTGGATATTAATGCCTTTCAACATTTCCTTAGAAAACTGGAATAAGGAAA
ATAAGAAAAACTTAATTCCACTTGAAAAGACTTACTTAGCTCCAATTCGTATCTATTCTGAGAAGGTAAAATCTCTTA
AAAAATTGAAAAAAGGAGCCACTATTGCAATTCCAAATGATGCAACAAATGGTAGCCGTGCATTGTATGTCCTTCAGT
CAGCAGGTTTAATCAAATTGAATGTTTCTGGTAAGAAGGTTGCAACAGTTGCTAATATCACATCTAATAAAAAGGATA
TTAATATTCAGGAGTTAGATGCGAGTCAAACACCACGTGCACTCAAAGATGTAGATGCAGCTATTATTAATAATACAT
ACATTGAGCAAGCTAATTTAAAACCTTCAGATGCTATCTTTGTTGAGAAATCAGATAAAAATTCAAAACAATGGATTA
ATATCATTGCGGGACGTAAAAATTGGAAAAAAGCAAAAGAACGCTAAAGCTATCCAAGCTATCTTGGATGCTTATCACA
CAGATGAAGTGAAAAAAAGTTATCAAAAGATACTTCAGCTGATATTCCACAATGG

### SEQ ID NO. 2909: SAG1641 FROM THE JM3190013 GBS TYPE VIII STRAIN

### SEQ ID NO. 2910: SAG1641 FROM THE M732 GBS TYPE III STRAIN



 ${\tt ATCAAGAAGTTTCAGCAAGCTCAACTTCAAGTAAAGTTGTTAAAGTTGGTGTTATGACC}$ 

SEQ2901

SEQESOI	ATCAAGAAGITICAGCAAGITCAAGITAAAGITGITAAAGITGITAIGACC
SEQ2902	ATCAAGAAGTTTCAGCAAGCTCAACTTCAAGTAAAGTTGTTAAAGTTGGTGTTATGACC
SEQ2903	ATCAAGAAGTTTCAGCAAGCTCAACTTCAAGTAAAGTTGTTAAAGTTGGTGTTATGACC
SEQ2904	ATCAAGAAGTTTCAGCAAGCTCAACTTCAAGTAAAGTTGTTAAAGTTGGTGTTATGACC
SEQ2905	ATCAAGAAGTTTCAGCAAGCTCAACTTCAAGTAAAGTTGTTAAAGTTGGTGTTATGACC
SEQ2906	
SEQ2907	AGTTTCAGCAAGCTCAACTTCAAGTAAAGTTGTTAAAGTTGGTGTTATGACC
SEQ2908	AAGAAGTTTCAGCAAGCTCAACTTCAAGTAAAGTTGTTTAAAGTTGGTGTTATGACC
SEQ2909	
<del>-</del>	AMENACA A COMMICA COA A COMMICA A COMICA A COMMICA A COM
SEQ2910	ATCAAGAAGTTTCAGCAAGCTCAACTTCAAGTAAAGTTGTTAAAGTTGGTGTTATGACC
SEQ2911	AGTTTCAGCAAGCTCAACTTCAAGTAAAGTTGTTAAAGTTGGTGTTATGACC
c=00004	
SEQ2901	TTTTCTGACACTGAAAAAGCACGTTGGGATAAAATTGAAAAGCTAGTAGGCGATAAAGCT
SEQ2902	TTTTCTGACACTGAAAAAGCACGTTGGGATAAAATTGAAAAGCTAGTAGGTGATAAAGCT
SEQ2903	TTTTCTGACACTGAAAAAGCACGTTGGGATAAAATTGAAAAGCTAGTAGGTGATAAAGCT
SEQ2904	TTTTCTGACACTGAAAAAGCACGTTGGGATAAAATTGAAAAGCTAGTAGGTGATAAAGCT
SEQ2905	TTTTCTGACACTGAAAAAGCACGTTGGGATAAAATTGAAAAGCTAGTAGGTGATAAAGCT
SEQ2906	TTTTCTGACACTGAAAAAGCACGTTGGGATAAAATTGAAAAGCTAGTAGGCGATAAAGCT
SEQ2907	TTTTCTGACACTGAAAAAGCACGTTGGGATAAAATTGAAAAGCTAGTAGGTGATAAAGCT
SEQ2908	TTTTCTGACACTGAAAAAGCACGTTGGGATAAAATTGAAAAGCTAGTAGGTGATAAAGCT
SEQ2909	TTTTCTGACACTGAAAAAGCACGTTGGGATAAAATTGAAAAGCTAGTAGGTGATAAAGCT
SEQ2910	TTTTCTGACACTGAAAAAGCACGTTGGGATAAAATTGAAAAGCTAGTAGGTGATAAAGCT
SEQ2911	TTTTCTGACACTGAAAAAGCACGTTGGGATAAAATTGAAAAGCTAGTAGGTGATAAAGCT
SEQ2901	AAAATCAAATTCACAGAATTTACAGATTATACACAACCAAATCAAGCGACAGCCAATAAG
SEQ2902	AAAATCAAATTTACAGAATTTACAGATTATACACAACCAAATCAAGCGACAGCCAATAAG
SEQ2903	AAAATCAAATTTACAGAATTTACAGATTATACACAACCAAATCAAGCGACAGCCAATAAG
SEQ2904	AAAATCAAATTTACAGAATTTACAGATTATACACAAATCAAGCGACAGCCAATAAG
SEQ2905	AAAATCAAATTTACAGAATTTACAGATTATACACAACCAAATCAAGCGACAGCCAATAAG
SEQ2906	AAAATCAAATTCACAGAATTTACAGATTATACACCAAATCAAGCGACAGCCAATAAG AAAATCAAATTCACAGAATTTACAGATTATACACCAAATCAAGCGACAGCCAATAAG
SEQ2907	AAAATCAAATTTACAGAATTTACAGATTATACACAACCAAATCAAGCGACAGCCAATAAG AAAATCAAATTTACAGAATTTACAGATTATACACAACCAAATCAAGCGACAGCCAATAAG
SEQ2908	
SEQ2909	AAAATCAAATTTACAGAATTTACAGATTATACACAAACCAAATCAAGCGACAGCCAATAAG
SEQ2909 SEQ2910	AAAATCAAATTTACAGAATTTACAGATTATACACAACCAAATCAAGCGACAGCCAATAAG
	AAAATCAAATTTACAGAATTTACAGATTATACACAACCAAATCAAGCGACAGCCAATAAG
SEQ2911	AAAATCAAATTTACAGAATTTACAGATTATACACAACCAAATCAAGCGACAGCCAATAAG
SEQ2901	GATGTGGATATTAATGCCTTTCAACATTACAATTTCTTAGAAAACTGGAATAAGGAAAAT
SEQ2902	
	GATGTGGATATTAATGCCTTTCAACATTACAATTTCTTAGAAAACTGGAATAAGGAAAAT
SEQ2903	GATGTGGATATTAATGCCTTTCAACATTACAATTTCTTAGAAAACTGGAATAAGGAAAAT
SEQ2904	GATGTGGATATTAATGCCTTTCAACATTACAATTTCTTAGAAAACTGGAATAAGGAAAAT
SEQ2905	GATGTGGATATTAATGCCTTTCAACATTACAATTTCTTAGAAAACTGGAATAAGGAAAAT
SEQ2906	GATGTGGATATTAATGCCTTTCAACATTACAATTTCTTAGAAAACTGGAATAAGGAAAAT
SEQ2907	GATGTGGATATTAATGCCTTTCAACATTACAATTTCTTAGAAAACTGGAATAAGGAAAAT
SEQ2908	GATGTGGATATTAATGCCTTTCAACATTACAATTTCTTAGAAAACTGGAATAAGGAAAAT
SEQ2909	GATGTGGATATTAATGCCTTTCAACATTACAATTTCTTAGAAAACTGGAATAAGGAAAAT
SEQ2910	GATGTGGATATTAATGCCTTTCAACATTACAATTTCTTAGAAAACTGGAATAAGGAAAAT
SEQ2911	GATGTGGATATTAATGCCTTTCAACATTACAATTTCTTAGAAAACTGGAATAAGGAAAAT
SEQ2901	AAGAAAACTTAATTCCACTTGAAAAGACTTACTTAGCCCCAATTCGTATCTATTCTGAG
SEQ2902	AAGAAAAACTTAATTCCACTTGAAAAGACTTACTTAGCTCCAATTCGTATCTATTCTGAG
SEQ2903	AAGAAAACTTAATTCCACTTGAAAAGACTTACTTAGCTCCAATTCGTATCTATTCTGAG
SEQ2904	AAGAAAACTTAATTCCACTTGAAAAGACTTACTTAGCTCCAATTCGTATCTATTCTGAG
SEQ2905	AAGAAAACTTAATTCCACTTGAAAAGACTTACTTAGCTCCAATTCGTATCTATTCTGAG
SEQ2906	AAGAAAACTTAATTCCACTTGAAAAGACTTACTTAGCCCCAATTCGTATCTATTCTGAG
SEQ2907	AAGAAAAACTTAATTCCACTTGAAAAGACTTACTTAGCTCCAATTCGTATCTATTCTGAG
SEQ2908	AAGAAAAACTTAATTCCACTTGAAAAGACTTACTTAGCTCCAATTCGTATCTATTCTGAG
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### Table 29: Comparative Sequences relating to SAG1641 (Yae-family pr tein)

22	
SEQZ909	AAGAAAAACTTAATTCCACTTGAAAAGACTTACTTAGCTCCAATTCGTATCTATTCTGAG
SEQ2910	AAGAAAAACTTAATTCCACTTGAAAAGACTTACTTAGCTCCAATTCGTATCTATTCTGAG
SEQ2911	AAGAAAACTTAATTCCACTTGAAAAGACTTACTTAGCTCCAATTCGTATCTATTCTGAG
SEQ2901	AAGGTAAAATCTCTTAAAAAATTGAAAAAAGGAGCCACTATTGCAATTCCAAATGATGCA
SEQ2902	AAGGTAAAATCTCTTAAAAAATTGAAAAAAGGAGCCACTATTGCAATTCCAAATGATGCA
SE02903	AAGGTAAAATCTCTTAAAAAATTGAAAAAAGGAGCCACTATTGCAATTCCAAATGATGCA
SEQ2904	AAGGTAAAATCTCTTAAAAAATTGAAAAAAGGAGCCACTATTGCAATTCCAAATGATGCA
SEQ2905	AAGGTAAAATCTCTTAAAAAATTGAAAAAAGGAGCCACTATTGCAATTCCAAATGATGCA
SEQ2906	AAGGTAAAATCTCTTAAAAAATTGAAAAAAGGAGCCACTATTGCAATTCCAAATGATGCA
SEQ2907	AAGGTAAAATCTCTTAAAAAATTGAAAAAAGGAGCCACTATTGCAATTCCAAATGATGCA
SEQ2908	AAGGTAAAATCTCTTAAAAAATTGAAAAAAGGAGCCACTATTGCAATTCCAAATGATGCA
SEQ2909	AAGGTAAAATCTCTTAAAAAATTGAAAAAAGGAGCCACTATTGCAATTCCAAATGATGCA
SEQ2910	AAGGTAAAATCTCTTAAAAAATTGAAAAAAGGAGCCACTATTGCAATTCCAAATGATGCA
SEQ2911	AAGGTAAAATCTCTTAAAAAATTGAAAAAAGGAGCCACTATTGCAATTCCAAATGATGCA
SEQ2901	ACAAATGGTAGCCGTGCATTGTATGTCCTTCAGTCAGCAGGTTTAATCAAATTGAATGTT
SEQ2902	ACAAATGGTAGCCGTGCATTGTATGTCCTTCAGTCAGCAGGTTTAATCAAATTGAATGTT
SEQ2903	ACAAATGGTAGCCGTGCATTGTATGTCCTTCAGTCAGCAGGTTTAATCAAATTGAATGTT
SEQ2904	ACAAATGGTAGCCGTGCATTGTATGTCCTTCAGTCAGCAGGTTTAATCAAATTGAATGTT
SEQ2905	ACAAATGGTAGCCGTGCATTGTATGTCCTTCAGTCAGCAGGTTTAATCAAATTGAATGTT
SEQ2906	ACAAATGGTAGCCGTGCATTGTATGTCCTTCAGTCAGCAGGTTTAATCAAATTGAATGTT
SEQ2907	ACAAATGGTAGCCGTGCATTGTATGTACTTCAGTCAGCAGGTTTAATCAAATTGAATGTT
SEQ2908	ACAAATGGTAGCCGTGCATTGTATGTCCTTCAGTCAGCAGGTTTAATCAAATTGAATGTT
SEQ2909	ACAAATGGTAGCCGTGCATTGTATGTCCTTCAGTCAGCAGGTTTAATCAAATTGAATGTT
SEQ2910	ACAAATGGTAGCCGTGCATTGTATGTCCTTCAGTCAGCAGGTTTAATCAAATTGAATGTT
SEQ2911	ACAAATGGTAGCCGTGCATTGTATGTCCTTCAGTCAGCAGGTTTAATCAAATTGAATGTT
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SEQ2901	TCTGGTAAGAAGGTTGCAACAGTTGCTAATATCACATCTAATAAAAAGGATATTAATATT
SEQ2902	TCTGGTAAGAAGGTTGCAACAGTTGCTAATATCACATCTAATAAAAAGGATATTAATATT
SEQ2903	TCTGGTAAGAAGGTTGCAACAGTTGCTAATATCACATCTAATAAAAAGGATATTAATATT
SEQ2904	TCTGGTAAGAAGGTTGCAACAGTTGCTAATATCACATCTAATAAAAAGGATATTAATATT
SEQ2905	TCTGGTAAGAAGGTTGCAACAGTTGCTAATATCACATCTAATAAAAAGGATATTAATATT
SEQ2906	TCTGGTAAGAAGGTTGCAACAGTTGCTAATATCACATCTAATAAAAAAGATATTAATATT
SEQ2907	TCTGGTAAGAAGGTTGCAACAGTTGCTAATATCACATCTAATAAAAAGGATATTAATATT
SEQ2908	TCTGGTAAGAAGGTTGCAACAGTTGCTAATATCACATCTAATAAAAAGGATATTAATATT
SEQ2909	TCTGGTAAGAAGGTTGCAACAGTTGCTAATATCACATCTAATAAAAAGGATATTAATATT
SEQ2910	TCTGGTAAGAAGGTTGCAACAGTTGCTAATATCACATCTAATAAAAAGGATATTAATATT
SEQ2911	TCTGGTAAGAAGGTTGCAACAGTTGCTAATATCACATCTAATAAAAAGGATATTAATATT
SEQ2901	CAGGAGTTAGATGCGAGTCAAACACCACGTGCACTCAAAGATGTAGATGCAGCTATTATT
SEQ2902	CAGGAGTTAGATGCGAGTCAAACACCACGTGCACTCAAAGATGTAGATGCAGCTATTATT
SEQ2903	CAGGAGTTAGATGCGAGTCAAACACCACGTGCACTCAAAGATGTAGATGCAGCTATTATT
SEQ2904	CAGGAGTTAGATGCCAGTCAAACACCACGTGCACTCAAAGATGTAGATGCAGCTATTATT
SEQ2905	CAGGAGTTAGATGCGAGTCAAACACCACGTGCACTCAAAGATGTAGATGCAGCTATTATT
SEQ2906	CAGGAGTTAGATGCGAGTCAAACACCACGTGCACTCAAAGATGTAGATGCAGCTATTATT
SEQ2907	CAGGAGTTAGATGCGAGTCAAACACCACGTGCACTCAAAGATGTAGATGCAGCTATTATT
SEQ2908 '	CAGGAGTTAGATGCGAGTCAAACACCACGTGCACTCAAAGATGTAGATGCAGCTATTATT
SEQ2909	CAGGAGTTAGATGCGAGTCAAACACCACGTGCACTCAAAGATGTAGATGCAGCTATTATT
SEQ2910	CAGGAGTTAGATGCGAGTCAAACACCACGTGCACTCAAAGATGTAGATGCAGCTATTATT
SEQ2911	CAGGAGTTAGATGCGAGTCAAACACCACGTGCACTCAAAGATGTAGATGCAGCTATTATT
CP02001	1 1 M 1 1 M 1 M 1 M 1 M 1 M 1 M 1 M 1 M
SEQ2901	AATAATACATACATTGAGCAAGCTAATTTAAAACCTTCAGATGCTATCTTTGTTGAGAAA
SEQ2902	AATAATACATACATTGAGCAAGCTAATTTAAAACCTTCAGATGCTATCTTTGTTGAGAAA
SEQ2903 SEO2904	AATAATACATACATTGAGCAAGCTAATTTAAAACCTTCAGATGCTATCTTTGTTGAGAAA
	AATAATACATACATTGAGCAAGCTAATTTAAAACCTTCAGATGCTATCTTTGTTGAGAAA
SEQ2905	AATAATACATACATTGAGCAAGCTAATTTAAAACCTTCAGATGCTATCTTTGTTGAGAAA
SEQ2906	AATAATACATACATTGAGCAAGCTAATTTAAAACCTTCAGATGCTATCTTTGTTGAGAAA
SEQ2907	AATAATACATACATTGAGCAAGCTAATTTAAAACCTTCAGATGCTATCTTTGTTGAGAAA
SEQ2908	AATAATACATTGAGCAAGCTAATTTAAAACCTTCAGATGCTATCTTTGTTGAGAAA
SEQ2909	AATAATACATACATTGAGCAAGCTAATTTAAAACCTTCAGATGCTATCTTTGTTGAGAAA
SEQ2910	AATAATACATTGAGCAAGCTAATTTAAAACCTTCAGATGCTATCTTTGTTGAGAAA
SEQ2911	AATAATACATACATTGAGCAAGCTAATTTAAAACCTTCAGATGCTATCTTTGTTGAGAAA

### Table 29: Comparative Sequences relating to SAG1641 (Yae Tamily pr tein)

SEQZ901	TCAGATAAAAATTCAAAACAATGGATTAATATCATTGCGGGACGTAAAAATTGGAAAAAG
SEQ2902	TCAGATAAAAATTCAAAACAATGGATTAATATCATTGCGGGACGTAAAAATTGGAAAAAG
SEQ2903	TCAGATAAAAATTCAAAACAATGGATTAATATCATTGCGGGACGTAAAAATTGGAAAAAG
SEQ2904	TCAGATAAAAATTCAAAACAATGGATTAATATCATTGCGGGACGTAAAAATTGGAAAAAG
SEQ2905	TCAGATAAAAATTCAAAACAATGGATTAATATCATTGCGGGACGTAAAAATTGGAAAAAG
SEQ2906	TCAGATAAAAATTCAAAACAATGGATTAATATCATTGCGGGACGTAAAAATTGGAAAAAG
SEQ2907	TCAGATAAAAATTCAAAACAATGGATTAATATCATTGCGGGACGTAAAAATTGGAAAAAG
SEQ2908	TCAGATAAAAATTCAAAACAATGGATTAATATCATTGCGGGACGTAAAAATTGGAAAAAG
SEQ2909	TCAGATAAAAATTCAAAACAATGGATTAATATCATTGCGGGACGTAAAAATTGGAAAAAG
SEQ2910	TCAGATAAAAATTCAAAACAATGGATTAATATCATTGCGGGACGTAAAAATTGGAAAAAG
SEQ2911	TCAGATAAAAATTCAAAACAATGGATTAATATCATTGCGGGACGTAAAAATTGGAAAAAG
SEQ2901	CAAAAGAACGCTAAAGCTATCCAAGCTATCTTGGATGCTTATCACACAGATGAAGTGAAA
SEQ2902	CAAAAGAACGCTAAAGCTATCCAAGCTATCTTGGATGCTTATCACACAGATGAAGTGAAA
SEQ2903	CAAAAGAACGCTAAAGCTATCCAAGCTATCTTGGATGCTTATCACACAGATGAAGTGAAA
SEQ2904	CAAAAGAACGCTAAAGCTATCCAAGCTATCTTGGATGCTTATCACACAGATGAAGTGAAA
SEQ2905	CAAAAGAACGCTAAAGCTATCCAAGCTATCTTGGATGCTTATCACACAGATGAAGTGAAA
SEQ2906	CAAAAGAACGCTAAAGCTATCCAAGCTATCTTGGATGCTTATCACACAGATGAAGTGAAA
SEQ2907	CAAAAGAACGCTAAAGCTATCCAAGCTATCTTGGATGCTTATCACACAGATGAAGTGAAA
SEQ2908	CAAAAGAACGCTAAAGCTATCCAAGCTATCTTGGATGCTTATCACACAGATGAAGTGAAA
SEQ2909	CAAAAGAACGCTAAAGCTATCCAAGCTATCTTGGATGCTTATCACACAGATGAAGTGAAA
SEQ2910	CAAAAGAACGCTAAAGCTATCCAAGCTATCTTGGATGCTTATCACACAGATGAAGTGAAA
SEQ2911	CAAAAGAACGCTAAAGCTATCCAAGCTATCTGGGATGCTTATCACACAGATGAAGTGAAA
SEQ2901	AAAGTTATCAAAGATACTTCAGCTGATATTCCACAATGGAACCCAGCTTTCTTGTACAA
SEQ2902	AAAGTTATCAAAGATACTTCAGCTGATATTCCACAATGG
SEQ2903	AAAGTTATCAAAGATACTTCAGCTGATATTCCAC
SEQ2904	AAAGTTATCAAAGATACTTCAGCTGATATTCCACAATGG
SEQ2905	AAAGTTATCAAAGATACTTCAGCTGATATTCCACAATGG
SEQ2906	AAAGTTATCAAAGATACTTCAGCTGATATTCCACAATGGAA
`SEQ2907	AAAGTTATCAAAGATACTTCAGCTGATATTCCACAATGG
SEQ2908	AAAGTTATCAAAGATACTTCAGCTGATATTCCACAATGG
SEQ2909	AAAGTTATCAAAGATACTTCAGCTGATATTCCACAATGG
SEQ2910	AAAGTTATCAAAGATAC
SEQ2911	AAAGTTATCAAAGATACTTCAGCTGATATTCCACAATGG

### >SEQ ID NO 2950: 35\_090 frame: 1

NQEVSASSTSSKVVKVGVMTFSDTEKARWDKIEKLVGDKAKIKFTEFTDYTQPNQATANK DVDINAFQHYNFLENWNKENKKNLIPLEKTYLAPIRIYSEKVKSLKKLKKGATIAIPNDA TNGSRALYVLQSAGLIKLNVSGKKVATVANITSNKKDINIQELDASQTPRALKDVDAAII NNTYIEQANLKPSDAIFVEKSDKNSKQWINIIAGRKNWKKQKNAKAIQAILDAYHTDEVK KVIKDTSADIPQWNPAFLY

### >SEQ ID NO 2951: 35\_1169NT frame: 3

QEVSASSTSKVVKVGVMTFSDTEKARWDKIEKLVGDKAKIKFTEFTDYTQPNQATANKD VDINAFQHYNFLENWNKENKKNLIPLEKTYLAPIRIYSEKVKSLKKLKKGATIAIPNDAT NGSRALYVLQSAGLIKLNVSGKKVATVANITSNKKDINIQELDASQTPRALKDVDAAIIN NTYIEQANLKPSDAIFVEKSDKNSKQWINIIAGRKNWKKQKNAKAIQAILDAYHTDEVKK VIKDTSADIPQW

### >SEQ ID NO 2952: 35\_18RS21 frame: 1

NQEVSASSTSSKVVKVGVMTFSDTEKARWDKIEKLVGDKAKIKFTEFTDYTQPNQATANK DVDINAFQHYNFLENWNKENKKNLIPLEKTYLAPIRIYSEKVKSLKKLKKGATIAIPNDA TNGSRALYVLQSAGLIKLNVSGKKVATVANITSNKKDINIQELDASQTPRALKDVDAAII NNTYIEQANLKPSDAIFVEKSDKNSKQWINIIAGRKNWKKQKNAKAIQAILDAYHTDEVK KVIKDTSADIP

#### >SEQ ID NO 2953:35 2603 frame: 1

NQEVSASSTSSKVVKVGVMTFSDTEKARWDKIEKLVGDKAKIKFTEFTDYTQPNQATANK DVDINAFQHYNFLENWNKENKKNLIPLEKTYLAPIRIYSEKVKSLKKLKKGATIAIPNDA TNGSRALYVLQSAGLIKLNVSGKKVATVANITSNKKDINIQELDASQTPRALKDVDAAII NNTYIEQANLKPSDAIFVEKSDKNSKQWINIIAGRKNWKKQKNAKAIQAILDAYHTDEVK KVIKDTSADIPQW

### Table 29: Comparative Sequences relating to SAG1641 (Yaec family pr tein)

### >SEQ ID NO 2954:35\_A909 frame: 1

NQEVSASSTSSKVVKVGVMTFSDTEKARWDKIEKLVGDKAKIKFTEFTDYTQPNQATANK DVDINAFQHYNFLENWNKENKKNLIPLEKTYLAPIRIYSEKVKSLKKLKKGATIAIPNDA TNGSRALYVLQSAGLIKLNVSGKKVATVANITSNKKDINIQELDASQTPRALKDVDAAII NNTYIEQANLKPSDAIFVEKSDKNSKQWINIIAGRKNWKKQKNAKAIQAILDAYHTDEVK KVIKDTSADIPQW

### >SEQ ID NO 2955:35\_CJB110 frame: 2

SKVVKVGVMTFSDTEKARWDKIEKLVGDKAKIKFTEFTDYTQPNQATANKDVDINAFQHY NFLENWNKENKKNLIPLEKTYLAPIRIYSEKVKSLKKLKKGATIAIPNDATNGSRALYVL QSAGLIKLNVSGKKVATVANITSNKKDINIQELDASQTPRALKDVDAAIINNTYIEQANL KPSDAIFVEKSDKNSKQWINIIAGRKNWKKQKNAKAIQAILDAYHTDEVKKVIKDTSADI PQW

### >SEQ ID NO 2956:35 COH1 frame: 2

VSASSTSSKVVKVGVMTFSDTEKARWDKIEKLVGDKAKIKFTEFTDYTQPNQATANKDVD INAFQHYNFLENWNKENKKNLIPLEKTYLAPIRIYSEKVKSLKKLKKGATIAIPNDATNG SRALYVLQSAGLIKLNVSGKKVATVANITSNKKDINIQELDASQTPRALKDVDAAIINNT YIEQANLKPSDAIFVEKSDKNSKQWINIIAGRKNWKKQKNAKAIQAILDAYHTDEVKKVI KDTSADIPOW

### >SEQ ID NO 2957:35\_H36B frame: 3

EVSÄSSTSSKVVKVGVMTFSDTEKARWDKIEKLVGDKAKIKFTEFTDYTQPNQATANKDV DINAFQHYNFLENWNKENKKNLIPLEKTYLAPIRIYSEKVKSLKKLKKGATIAIPNDATN GSRALYVLQSAGLIKLNVSGKKVATVANITSNKKDINIQELDASQTPRALKDVDAAIINN TYIEQANLKPSDAIFVEKSDKNSKQWINIIAGRKNWKKQKNAKAIQAILDAYHTDEVKKV IKDTSADIPOW

#### >SEQ ID NO 2958:35 JM9130013 frame: 2

SASSTSSKVVKVGVMTFSDTEKARWDKIEKLVGDKAKIKFTEFTDYTQPNQATANKDVDI NAFQHYNFLENWNKENKKNLIPLEKTYLAPIRIYSEKVKSLKKLKKGATIAIPNDATNGS RALYVLQSAGLIKLNVSGKKVATVANITSNKKDINIQELDASQTPRALKDVDAAIINNTY IEQANLKPSDAIFVEKSDKNSKQWINIIAGRKNWKKQKNAKAIQAILDAYHTDEVKKVIK DTSADIPQW

### >SEQ ID NO 2959:35\_M732 frame: 1

NQEVSASSTSSKVVKVGVMTFSDTEKARWDKIEKLVGDKAKIKFTEFTDYTQPNQATANK DVDINAFQHYNFLENWNKENKKNLIPLEKTYLAPIRIYSEKVKSLKKLKKGATIAIPNDA TNGSRALYVLQSAGLIKLNVSGKKVATVANITSNKKDINIQELDASQTPRALKDVDAAII NNTYIEQANLKPSDAIFVEKSDKNSKQWINIIAGRKNWKKQKNAKAIQAILDAYHTDEVK KVIKD

### >SEQ ID NO 2960:35 M781 frame: 2

VSASSTSSKVVKVGVMTFSDTEKARWDKIEKLVGDKAKIKFTEFTDYTQPNQATANKDVD INAFQHYNFLENWNKENKKNLIPLEKTYLAPIRIYSEKVKSLKKLKKGATIAIPNDATNG SRALYVLQSAGLIKLNVSGKKVATVANITSNKKDINIQELDASQTPRALKDVDAAIINNT YIEQANLKPSDAIFVEKSDKNSKQWINIIAGRKNWKKQKNAKAIQAIWDAYHTDEVKKVI KDTSADIPQW

SEQ2950	QEVSASSTSSKVVKVGVMTFSDTEKARWDKIEKLVGDKAKIKFTEFTDYTQPNQATANK
SEQ2951	QEVSASSTSSKVVKVGVMTFSDTEKARWDKIEKLVGDKAKIKFTEFTDYTQPNQATANK
SEQ2952	OF UCA COMOCULARIZATION DE DESIGNATURA DE LA COMOCULARIZATION DEL COMOCULARIZATION DE LA CO
	QEVSASSTSSKVVKVGVMTFSDTEKARWDKIEKLVGDKAKIKFTEFTDYTQPNQATANK
SEQ2953	QEVSASSTSSKVVKVGVMTFSDTEKARWDKIEKLVGDKAKIKFTEFTDYTQPNQATANK
SEQ2954	QEVSASSTSSKVVKVGVMTFSDTEKARWDKIEKLVGDKAKIKFTEFTDYTQPNQATANK
SEQ2955	SKVVKVGVMTFSDTEKARWDKIEKLVGDKAKIKFTEFTDYTQPNQATANK
SEQ2956	VSASSTSSKVVKVGVMTFSDTEKARWDKIEKLVGDKAKIKFTEFTDYTQPNQATANK
SEQ2957	-EVSASSTSSKVVKVGVMTFSDTEKARWDKIEKLVGDKAKIKFTEFTDYTQPNQATANK
SEQ2958	CA COMOCIATIVA DE LE COMPANIA DE LA COMPONIA DEL COMPONIA DEL COMPONIA DE LA COMPONIA DE LA COMPONIA DEL COMPONICA DEL COMPONIA DEL COMPONICA DEL COMPONIA DEL COMPONIO DEL CO
<del></del>	SASSTSSKVVKVGVMTFSDTEKARWDKIEKLVGDKAKIKFTEFTDYTQPNQATANK
SEQ2959	QEVSASSTSSKVVKVGVMTFSDTEKARWDKIEKLVGDKAKIKFTEFTDYTQPNQATANK
SEQ2960	VSASSTSSKVVKVGVMTFSDTEKARWDKIEKLVGDKAKIKFTEFTDYTQPNQATANK
SEQ2950	DUDTNA POHVNET PNUMVENVVNI TDI EVENUS SETE TUONG WOLLDON WAR
<del>-</del> .	DVDINAFQHYNFLENWNKENKKNLIPLEKTYLAPIRIYSEKVKSLKKLKKGATIAIPNDA
SEQ2951	DVDINAFQHYNFLENWNKENKKNLIPLEKTYLAPIRIYSEKVKSLKKLKKGATIAIPNDA
SEQ2952	DVDINAFQHYNFLENWNKENKKNLIPLEKTYLAPIRIYSEKVKSLKKLKKGATIAIPNDA
SEQ2953	DVDINAFQHYNFLENWNKENKKNLIPLEKTYLAPIRIYSEKVKSLKKLKKGATIAIPNDA
SEQ2954	DVDINAFQHYNFLENWNKENKKNLIPLEKTYLAPIRIYSEKVKSLKKLKKGATIAIPNDA
SEQ2955	DUDTIN POUNTY BURNISHED FOR THE TRI I SERVASERERRORI I ALFINDA
	DVDINAFQHYNFLENWNKENKKNLIPLEKTYLAPIRIYSEKVKSLKKLKKGATIAIPNDA
SEQ2956	DVDINAFQHYNFLENWNKENKKNLIPLEKTYLAPIRIYSEKVKSLKKLKKGATIAIPNDA
SEQ2957	DVDINAFQHYNFLENWNKENKKNLIPLEKTYLAPIRIYSEKVKSLKKLKKGATIAIPNDA
SEQ2958	DVDINAFQHYNFLENWNKENKKNLIPLEKTYLAPIRIYSEKVKSLKKLKKGATIAIPNDA
	2. 52. W. WILL DOWNLOAD AND PLEKTYLAPIRTY SEKVKSLKKLKKGATTALPNDA



SE02959	DVDINAFOHYNFLENWNKENKKNLIPLEKTYLAPIRIYSEKVKSLKKLKKGATIAIPNDA
SEQ2960	DVDINAFQHYNFLENWNKENKKNLIPLEKTYLAPIRIYSEKVKSLKKLKKGATIAIPNDA
SEQ2950	TNGSRALYVLQSAGLIKLNVSGKKVATVANITSNKKDINIQELDASOTPRALKDVDAAII
SEQ2951	TNGSRALYVLQSAGLIKLNVSGKKVATVANITSNKKDINIQELDASQTPRALKDVDAAII
SEQ2952	TNGSRALYVLQSAGLIKLNVSGKKVATVANITSNKKDINIQELDASQTPRALKDVDAAII
SEQ2953	TNGSRALYVLQSAGLIKLNVSGKKVATVANITSNKKDINIQELDASQTPRALKDVDAAII
SEQ2954	TNGSRALYVLQSAGLIKLNVSGKKVATVANITSNKKDINIQELDASQTPRALKDVDAAII
SEQ2955	TNGSRALYVLQSAGLIKLNVSGKKVATVANITSNKKDINIQELDASQTPRALKDVDAAII
SEQ2956	TNGSRALYVLQSAGLIKLNVSGKKVATVANITSNKKDINIQELDASQTPRALKDVDAAII
SEQ2957	TNGSRALYVLQSAGLIKLNVSGKKVATVANITSNKKDINIQELDASQTPRALKDVDAAII
SEQ2958	TNGSRALYVLQSAGLIKLNVSGKKVATVANITSNKKDINIQELDASQTPRALKDVDAAII
SEQ2959	TNGSRALYVLQSAGLIKLNVSGKKVATVANITSNKKDINIQELDASQTPRALKDVDAAII
SEQ2960	TNGSRALYVLQSAGLIKLNVSGKKVATVANITSNKKDINIQELDASQTPRALKDVDAAII
SEQ2950	NNTYIEQANLKPSDAIFVEKSDKNSKQWINIIAGRKNWKKQKNAKAIQAILDAYHTDEVK
SEQ2951	NNTYIEQANLKPSDAIFVEKSDKNSKQWINIIAGRKNWKKQKNAKAIQAILDAYHTDEVK
SEQ2952	NNTYIEQANLKPSDAIFVEKSDKNSKQWINIIAGRKNWKKQKNAKAIQAILDAYHTDEVK
SEQ2953	NNTYIEQANLKPSDAIFVEKSDKNSKQWINIIAGRKNWKKQKNAKAIQAILDAYHTDEVK
SEQ2954	NNTYIEQANLKPSDAIFVEKSDKNSKQWINIIAGRKNWKKQKNAKAIQAILDAYHTDEVK
SEQ2955	NNTYIEQANLKPSDAIFVEKSDKNSKQWINIIAGRKNWKKQKNAKAIQAILDAYHTDEVK
SEQ2956	NNTYIEQANLKPSDAIFVEKSDKNSKQWINIIAGRKNWKKQKNAKAIQAILDAYHTDEVK
SEQ2957	NNTYIEQANLKPSDAIFVEKSDKNSKQWINIIAGRKNWKKQKNAKAIQAILDAYHTDEVK
SEQ2958	NNTYIEQANLKPSDAIFVEKSDKNSKQWINIIAGRKNWKKQKNAKAIQAILDAYHTDEVK
SEQ2959	NNTYIEQANLKPSDAIFVEKSDKNSKQWINIIAGRKNWKKQKNAKAIQAILDAYHTDEVK
SEQ2960	NNTYIEQANLKPSDAIFVEKSDKNSKQWINIIAGRKNWKKQKNAKAIQAIWDAYHTDEVK
SEQ2950	KVIKDTSADIPOWNPAFLY
SEQ2951	KVIKDTSADIPQW
SEQ2952	KVIKDTSADIP
SEQ2953	KVIKDTSADIPOW
SEQ2954	KVIKDTSADIPOW
SEQ2955	KVIKDTSADIPOW
SEQ2956	KVIKDTSADIPOW
SEQ2957	KVIKDTSADIPQW
SEQ2958	KVIKDTSADIPQW
SEQ2959	KVIKD
SEQ2960	KVIKDTSADIPQW

## Table 30: Comparative Sequences relating to SAG2147 (protein f uknown function / lipoprotein, putative)

### SEQ ID NO. 3001: SAG2147 FROM THE 1169NT1 GBS TYPE V STRAIN (REVERSE COMPLEMENT)

AAAAGTTCACAAGTTACTGCTGAATCTTTGTCAAAAGCAGATAAAGTTCGCGTAGCC
AAAAAATCAAAAATGACTAAGGCGACATCTAAATCAAAAGTAGAAGATGTAAAACAGGCT
CCAAAACCTTCTCAGGCATCTAATGAAGTCCCAAAATCAAGTTCTCAATCTACAGAAGCT
AATTCTCAGCAACAAGTTACTGCGAGTGAAGAGGCGGCTGTAGAACAAGCAGTTGTAACA
GAAAATACCCCTGCTACCAGTCAGGCACAACAAACTTATGCTGTTACTGAGACAACTTAC
AAACCTGCTCAACACCAGACAAGTGGCCAAGTATTGAGCAATGGAAATACTGCAGGGGCG
GTCGGATCTGCTGCAGCACAAATGGCTGCTGCAACAGGAGTCCCTCAGTCTACTTGG
GAACATATTATTGCCCGTGAATCAAATGGTAATCCTAATGTTGCTAATGCCTCAGGAGCT
TCAGGACTTTTCCAAACGATGCCAGGTTGGGGTTCAACAGCTTCAGGATCAAGTT
AATTCAGCTATTAAAGCTTATCGTGCTCAAAGGTTTATCAGCTTGGGGTTAC

### SEQ ID NO. 3002: SAG2147 FROM THE 18RS21 GBS TYPE II STRAIN (REVERSE COMPLEMENT)

AAAAGTTCACAAGTTACTGCTGAATCTTTGTCAAAAGCAGATAAAGTTC
GCGTAGCCAAAAAATCAAAAATGACTAAGGCGACATCTAAATCAAAAGTAGAAGATGTAA
AACAGGCTCCAAAACCTTCTCAGGCATCTAATGAAGCCCCAAAATCAAGTTCTCAATCTA
CAGAAGCTAATTCTCAGCAACAAGTTACTGCGAGTGAAGAGGCAGCTGTAGAACAAGCAG
TTGTAACAGAAAACACCCCTGCTACCAGTCAGGCACAACAAGCTTATGCTGTTACTGAGA
CAACTTATAGACCTGCTCAACACCAGACGAGTGGCCAAGTATTGAGTAATGGAAATACTG
CAGGGGCTATTGGCTCAGCAGCTGCAGCACAAATGGCTGCTGCAACAGGAGTCCCTCAGT
CTACTTGGGAACATATTATTGCCCGTGAATCAAATGGTAATCCTAATGTTGCTAATGCCT
CAGGAGCTTCAGGACTTTTCCAAACGATGCCAGGTTGGGGTTCAACAGCTACAGTTCAGG
ATCAAGTTAATTCAGCTATTAAAGCTTATCGTGCTCCAAGGTTTATCAGCTTTGGGGTTAC

### SEQ ID NO. 3003: SAG2147 FROM THE 2603 V/R GBS TYPE V STRAIN (REVERSE COMPLEMENT)

AAAAGTTCACAAGTTACTACTGAATCTTTGTCAAAAGCAGATAAAGT
TCGCGTAGCCAAAAAATCAAAAATGACTAAGGCGACATCTAAATCAAAAGTAGAAGATGT
AAAACAGGCTCCAAAACCTTCTCAGGCATCTAATGAAGCCCCCAAAATCAAGTTCTCAATC
TACAGAAGCTAATTCTCAGCAACAAGTTACTGCGAGTGAAGAGGCAGCTGTAGAACAAGC
AGTTGTAACAGAAAACACCCCTGCTACCAGTCAGGCACAACAAGCTTATGCTGTTACTGA
GACAACTTATAGACCTGCTCAACACCAGACGAGTGGCCAAGTATTGAGTAATGGAAATAC
TGCAGGGGCTATTGGCTCAGCAGCTGCAGCACAAATGGCTGCTAACAGGAGTCCCTCA
GTCTACTTGGGAACATATTATTGCCCGTGAATCAAATGGTAATCCTAATGTTGCTAATGC
CTCAGGAGCTTCAGGACTTTTCCAAACGATGCCAGGTTCAACAGCTACAGTTCA
GGATCAAGTTAATTCAGCTATTAAAGCTTATCGTGCTCAAGGTTTATCAGCTTGGGGTTA

### SEQ ID NO. 3004: SAG2147 FROM THE 090 GBS TYPE IA STRAIN (REVERSE COMPLEMENT)

TAGCCAAAAATCAAAAATGATTAAGGCGACATCTAAATCAAAAGTAGAAGATGTAAAAC
AGGCTCCAAAACCTTCTCAGGCATCTAATGAAGCCCCCAAAATCAAGTTCTCAATCTACAG
AAGCTAATTCTCAGCAACAAGTTACTGCGAGTGAAGAGGCAGCTGTAGAACAAGCAGTTG
TAACAGAAAACACCCCTGCTACCAGTCAGGCACAACAAGCTTATGCTGTTACTGAGACAA
CTTATAGACCTGCTCAACACCAGACGAGTGGCCCAAGTATTGAGTAATGGAAATACTGCAG
GGGCTATTGGCTCAGCAGCTGCAGCACAAATGGCTGCTGCAACAGGAGTCCCTCAGTCTA
CTTGGGAACATATTATTGCCCGTGAATCAAATGGTAATCCTAATGTTGCTAATGCCTCAG
GAGCTTCAGGACTTTCCAAACGATGCCAGGTTGGGGTTCAACAGCTACAGTTCAGGA

### SEQ ID NO. 3005: SAG2147 FROM THE A909 GBS TYPE In STRAIN (REVERSE COMPLEMENT)

AAGGCGACATCTAAATCAAAAGTAGAAGATGTAAAACAGGCTCCAAAACCTTCTCAGGCA
TCTAATGAAGCCCCAAAATCAAGTTCTCAATCTACAGAAGCTAATTCTCAGCAACAAGTT
ACTGCGAGTGAAGAGGCAGCTGTAGAACAAGCAGTTGTAACAGAAAACACCCCTGCTACC
AGTCAGGCACAACAAGCTTATGCTGTTACTGAGACAACTTATAGACCTGCTCAACACCAG
ACAAGTGGCCAAGTATTGAGTAATGGAAATACTGCAGGGGCTATTGGCTCAGCAGCTGCA
GCACAAATGGCTGCTAACAGGAGTCCCTCAGTCTACTTGGGAACATATTATTGCCCGT
GAATCAAATGGTAATCCTAATGTTGCTAATGCCTCAGGAGCTTCAGGACTTTTCCAAACG
ATGCCAGGTTGGGGTTCAACAGCTACAGTTCAGAATCAAGTTAATTCAGCTATTAAAGCT
TATCGTGCTCAAGGTTTATCA

## Table 30: Comparative Sequences relating t SAG2147 (pr tein of uknown functi n / lipoprotein, putative)

### SEQ ID NO. 3006: SAG2147 FROM THE CJB110 GBS NONTYPEABLE STRAIN (REVERSE COMPLEMENT)

AATCTTTGTCAAAAGCAGATAAAGTTCGCGTAGCCAAAAAATCAAAAATGACTAAGGCGA
CATCTAAATCAAAAGTAGAAGATGTAAAACAGGCTCCAAAACCTTCTCAGGCATCTAATG
AAGCCCCAAAATCAAGTTCTCAATCTACAGAAGCTAATTCTCAGCAACAAGTTACTGCGA
GTGAAGAGGCAGCTGTAGAACAAGCAGTTGTAACAGAAAACACCCCTGCTACCAGTCAGG
CACAACAAGCTTATGCTGTTACTGAGACAACTTATAGACCTGCTCAACACCAGACGAGTG
GCCAAGTATTGAGTAATGGAAATACTGCAGGGGCTATTGGCTCAGCAGCTGCAGCACAAA
TGGCTGCTGCAACAGGAGTCCCTCAGTCTACTTGGGAACATATTATTGCCCGTGAATCAA
ATGGTAATCCTAATGTTGCTAATGCCTCAGGAGCTTCAGGACTTTTCCAAACGATGCCAG
GTTGGGGTTCAACAGCTACAGTTCAGGATCAAGTTAATTCAGCTATTAAAGCTTATCGTG
CTCAAGGTTTATCAGCTTGGGGTTAC

### SEQ ID NO. 3007: SAG2147 FROM THE COH1 GBS TYPE III STRAIN (REVERSE COMPLEMENT)

AAAAGTTCACAAGTTACTACTGAATCTTTGTCAAAAGCAGATAA
AGTTCGCGTAGCCAAAAAATCAAAAATGACTAAGGCGACATCTAAATCAAAAGTAGAAGA
TGTAAAACAGGCTCCAAAACCTTCTCAGGCATCTAATGAAGCCCCAAAATCAAGTTCTCA
ATCTACAGAAGCTAATTCTCAGCAACAAGTTACTGCGAGTGAAGAGGCGGCTGTAGAACA
AGCAGTTGTAACAGAAAATACCCCTGCTACCAGTCAGGCACAACAAACTTATGCTGTTAC
TGAGACAACTTACAAACCTGCTCAACACCAGACAAGTGGCCAAGTATTGAGCAATGGAAA
TACTGCAGGGGCGGTCGGATCTGCTGCTGCAGCACAAATGGCTGCTGCAACAGGAGTCCC
TCAGTCTACTTGGGAACATATTATTGCCCGTGAATCAAATGGTAATCCTAATGTTGCTAA
TGCCTCAGGAGCTTCAGGACTTTTCCAAACGATGCCAGGTTGGGGTTCAACAGCTACAGT
TCAGGATCAAGTTAATTCAGCTATTAAAGCTTATCGTGCTCAAGGTTTATCAGCTTGGGG
TTAC

### SEQ ID NO. 3008: SAG2147 FROM THE H36b GBS TYPE Ib STRAIN (REVERSE COMPLEMENT)

AAAAGTTCACAAGTTACTACTGAATCTTTGTCAAAAGC

AGATAAAGTTCGCGTAGCCAAAAAATCAAAAATGACTAAGGCGACATCTAAATCAAAAGT AGAAGATGTAAAAACAGCCCCAAAAACCTTCTCAGGCATCTAATGAAGCCCCAAAATCAAG TTCTCAATCTAAAAACCCCCAAAATCAAG TTCTCAATCTACAGAAGCCAACAAGTTACTGCGAGTGAAGAGGCAGCTGT AGAACAAGCAGTTGTAACAGAAAACACCCCTGCTACCAGTCAGGCACAACAAGCTTATGC TGTTACTGAGACAACTTATAGACCTGCTCAACACCAGACAAGTGGCCAAGTATTGAGTAA TGGAAATACTGCAGGGGCTATTGGCTCAGCAGCTGCAGCACAAATGGCTGCAACAGG AGTCCCTCAGTCTACTTGGGAACATTATTTTCCCCGTGAATCAAATGGTAATCCTAATGT TGCTAATGCCTCAGGAGCTTCAGGACTTTCCAAACGATGCCAGGTTGGGGTTCAACAGC TACAGTTCAGGATCAAATTAATTCAGCTATTAAAGCTT

### SEQ ID NO. 3009: SAG2147 FROM THE M732 GBS TYPE III STRAIN (REVERSE COMPLEMENT)

AAAAGTTCACAAGTTACTACTGAATCTTTGTCAAAAGCAGATAAAGTTCGCGTAGC
CAAAAAATCAAAAATGACTAAGGCGACATCTAAATCAAAAGTAGAAGATGTAAAACAGGC
TCCAAAACCTTCTCAGGCATCTAATGAAGCCCCCAAAATCAAGTTCTCAATCTACAGAAGC
TAATTCTCAGCAACAAGTTACTGCGAGTGAAGAGGCGGCTGTAGAACAAGCAGTTGTAAC
AGAAAATACCCCTGCTACCAGTCAGGCACAACAAACTTATGCTGTTACTGAGACAACTTA
CAAACCTGCTCAACACACAAAGTGGCCAAGTATTGAGCAATGGAAATACTGCAGGGGC
GGTCGGATCTGCTGCAGCACAAATGGCTGCTGCAACAGGAGTCCCTCAGTCTACTTG
GGAACATATTATTGCCCGTGAATCAAATGGTAATCCTAATGTTGCTAATGCCTCAGGAGC
TTCAGGACTTTTCCAAACGATGCCAGGTTGGGGTTCAACAGCTACAGTTCAGGATCAAGT
TAATTCAGCTATTAAAGCTTATCGTGCTCAAGGTTTATCAGCTTGGGGTTA



# Table 30: Comparativ Sequences relating t SAG2147 (protein of uknown function / lipoprotein, putative)

SEQ ID NO. 3010: SAG2147 FROM THE M781 GBS TYPE III STRAIN (REVERSE COMPLEMENT)

SEQ3001	~
SEQ3002	~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~
SEQ3003	
SEQ3004	
SEQ3005	AGGCGACATCTAAATCAAAAGTAGAAGATGTAAAACAGGCTCCAAAACCTTCTCAGGCA
SEQ3007	
SEQ3008	
SEQ3009	
SEQ3010	
SEQ3001	
SEQ3002	
SEQ3003	
SEQ3004	
SEQ3005	CTAATGAAGCCCCAAAATCAAGTTCTCAATCTACAGAAGCTAATTCTCAGCAACAAGTT
SEQ3007	
SEQ3008	~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~
SEQ3009	***************************************
SEQ3010	
SEQ3001	
SEQ3002	
SEQ3003	
SEQ3004	
SEQ3005	CTGCGAGTGAAGAGGCAGCTGTAGAACAAGCAGTTGTAACAGAAAACACCCCTGCTACC
SEQ3007	
SEQ3008	
SEQ3009	
SEQ3010	***********************************
SEQ3001	
SEQ3002	
SEQ3003	
SEQ3004	
SEQ3005	GTCAGGCACAACAAGCTTATGCTGTTACTGAGACAACTTATAGACCTGCTCAACACCAG
SEQ3007	GICHOGGAGEROGIA INIGOTO I ACCOMONCACE I MINORCEIGEI CAMCACCAG
SEQ3008	H-04-04-04-04-04-04-04-04-04-04-04-04-04-
SEQ3009	
SEQ3010	
5525010	
SEQ3001	
SEQ3002	
SEQ3003	4-4
SEQ3004	
SEQ3005	CAAGTGGCCAAGTATTGAGTAATGGAAATACTGCAGGGGCTATTGGCTCAGCAGCTGCA
SEQ3007	CARGIGGCCAAGIAIIGAAAATACTGCAGGGGCTATTGGCTCAGCAGCTGCA
SEQ3008	
SEQ3009	
SEQ3010	=======================================
SECOUTA	

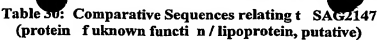
# Table 30: C mparative Sequences relating t SAG2147 (protein of uknown function / lipopr tein, putative)

SEQ3001	
SEQ3002	
SEQ3003	
_	
SEQ3004	
SEQ3005	CACAAATGGCTGCTACAGGAGTCCCTCAGTCTACTTGGGAACATATTATTGCCCGT
SEQ3007	
SEQ3008	
SEQ3009	
SEQ3010	
2280070	
GE02001	
SEQ3001	
SEQ3002	
SEQ3003	
SEQ3004	*
SEQ3005	AATCAAATGGTAATCCTAATGTTGCTAATGCCTCAGGAGCTTCAGGACTTTTCCAAACG
SEQ3007	
SEQ3008	
SEQ3009	***************************************
SEQ3010	***************************************
SEQ3001	+
SEQ3002	
SEQ3003	
SEQ3004	
SEQ3005	TGCCAGGTTGGGGTTCAACAGCTACAGTTCAGAATCAAGTTAATTCAGCTATTAAAGCT
SEQ3003 SEQ3007	TGCCAGGTTGGGGTTCAACAGCTACAGTTCAGAATCAAGTTAATTCAGCTATTAAAGCT
-	
SEØ3008	
SEQ3009	
SEQ3010	
SEQ3001	AAAGTTCACAAGTTACTACTGAATCTTTGTCAAAAGCAGATAAAGTTCGCGTAGCCAAA
SEQ3002	AAAGTTCACAAGTTACTGAATCTTTGTCAAAAGCAGATAAAGTTCGCGTAGCCAAA
SEQ3003	AAAGTTCACAAGTTACTGAATCTTTGTCAAAAGCAGATAAAGTTCGCGTAGCCAAA
SEQ3004	TAGCCAAA
SEQ3005	
	ATCGTGCTCAAGGTTTATCASAATCTTTGTCAAAAGCAGATAAAGTTCGCGTAGCCAAA
SEQ3007	AAAGTTCACAAGTTACTGAATCTTTGTCAAAAGCAGATAAAGTTCGCGTAGCCAAA
SEQ3008	AAAGTTCACAAGTTACTGAATCTTTGTCAAAAGCAGATAAAGTTCGCGTAGCCAAA
SEQ3009	AAAGTTCACAAGTTACTACTGAATCTTTGTCAAAAGCAGATAAAGTTCGCGTAGCCAAA
SEQ3010	GTAACCCCAAGCTGATAAACCTTGAGCACGATAAGCTTTAATAGCTGAA
SEQ3001	AATCAAAAATGACTAAGGCGACATCTAAATCAAAAGTAGAAGATGTAAAACAGGCTCCA
SEQ3002	AATCAAAAATGACTAAGGCGACATCTAAATCAAAAGTAGAAGATGTAAAACAGGCTCCA
SEQ3003	AATCAAAAATGACTAAGGCGACATCTAAATCAAAAGTAGAAGATGTAAAACAGGCTCCA
SEQ3004	
	AATCAAAAATGATTAAGGCGACATCTAAATCAAAAGTAGAAGATGTAAAACAGGCTCCA
SEQ3005	AATCAAAAATGACTAAGGCGACATCTAAATCAAAAGTAGAAGATGTAAAACAGGCTCCA
SEQ3007	AATCAAAAATGACTAAGGCGACATCTAAATCAAAAGTAGAAGATGTAAAACAGGCTCCA
SEQ3008	AATCAAAAATGACTAAGGCGACATCTAAATCAAAAGTAGAAGATGTAAAACAGGCTCCA
SEQ3009	AATCAAAAATGACTAAGGCGACATCTAAATCAAAAGTAGAAGATGTAAAACAGGCTCCA
SEQ3010	TAACTTGATCCTGAACTGTAGCTGTTGAACCCCAACCTGGCATCGTTTGGAAAAGTCCT
SEQ3001	AACCTTCTCAGGCATCTAATGAAGTCCCAAAATCAAGTTCTCAATCTACAGAAGCTAAT
SEQ3002	AACCTTCTCAGGCATCTAATGAAGCCCCAAAATCAAGTTCTCAATCTACAGAAGCTAAT AACCTTCTCAGGCATCTAATGAAGCCCCCAAAATCAAGTTCTCAATCTACAGAAGCTAAT
	AMOUTTOT CAGGEAT CTAATGAAGCCCCAAAATCAAGTTCTCAATCTACAGAAGCTAAT
SEQ3003	AACCTTCTCAGGCATCTAATGAAGCCCCAAAATCAAGTTCTCAATCTACAGAAGCTAAT
SEQ3004	AACCTTCTCAGGCATCTAATGAAGCCCCAAAATCAAGTTCTCAATCTACAGAAGCTAAT
SEQ3005	AACCTTCTCAGGCATCTAATGAAGCCCCAAAATCAAGTTCTCAATCTACAGAAGCTAAT
SEQ3007	AACCTTCTCAGGCATCTAATGAAGCCCCAAAATCAAGTTCTCAATCTACAGAAGCTAAT
SEQ3008	AACCTTCTCAGGCATCTAATGAAGCCCCAAAATCAAGTTCTCAATCTACAGAAGCTAAT
SEQ3009	AACCTTCTCAGGCATCTAATGAAGCCCCCAAAATCAAGTTCTCAATCTACAGAAGCTAAT
SEQ3010	AAGCTCCTGAGGCATTAGCAACATTAGGATTAC-CATTTGATTCACGGGCAATAAT
25010	AMSCICCIGAGGCAIIAGCAACATTAGGATTAC-CATTTGATTCACGGGCAATAAT
000001	MOMO3 COL 3 C3 3 CMM3 CMC COL 3 CMC CA
SEQ3001	TCTCAGCAACAAGTTACTGCGAGTGAAGAGGGGGGCTGTAGAACAAGCAGTTGTAACAGA
SEQ3002	TCTCAGCAACAAGTTACTGCGAGTGAAGAGGCAGCTGTAGAACAAGCAGTTGTAACAGA
SEQ3003	TCTCAGCAACAAGTTACTGCGAGTGAAGAGGCAGCTGTAGAACAAGCAGTTGTAACAGA
SEQ3004	TCTCAGCAACAAGTTACTGCGAGTGAAGAGGCAGCTGTAGAACAAGCAGTTGTAACAGA
SEQ3005	TCTCAGCAACAAGTTACTGCGAGTGAAGAGGCAGCTGTAGAACAAGCAGTTGTAACAGA
SEQ3007	TCTCAGCAACAAGTTACTGCGAGTGAAGAGGCGGCTGTAGAACAAGCAGTTGTAACAGA
SEQ3008	TOTOLOGIA A CARCETTO TO COLOR CONTROL CARCETTO TARCALA CA
**	TCTCAGCAACAAGTTACTGCGAGTGAAGAGGCAGCTGTAGAACAAGCAGTTGTAACAGA
SEQ3009	TCTCAGCAACAAGTTACTGCGAGTGAAGAGGCGGCTGTAGAACAAGCAGTTGTAACAGA
SEQ3010	TGTTCCCAAGTAGACTGAGGGACTCCTGTTGCAGCAGCCATTTGTGCTGCAGCAGCAGA



# Table 50: Comparative Sequences relating t SAG2147 (protein of uknown function / lipoprotein, putative)

SEQ3001	Aaatacccctgctaccagtcaggcacaacaaacttatgctgttactgagacaactta
SEQ3002	AAACACCCCTGCTACCAGTCAGGCACAACAAGCTTATGCTGTTACTGAGACAACTTA
SE03003	AAACACCCCTGCTACCAGTCAGGCACAACAAGCTTATGCTGTTACTGAGACAACTTA
SEQ3004	AAACACCCCTGCTACCAGTCAGGCACAACAAGCTTATGCTGTTACTGAGACAACTTA
SEQ3005	AAACACCCCTGCTACCAGTCAGGCACAACAAGCTTATGCTGTTACTGAGACAACTTA
SEQ3007	AAATACCCCTGCTACCAGTCAGGCACAACATTATGCTGTTACTGAGACAACTTA
SEQ3008	ANALOGO TO CALONDO CALONDO CALONDO CONTROL CON
SEQ3009	AAACACCCCTGCTACCAGTCAGGCACAACAAGCTTATGCTGTTACTGAGACAACTTA
SEQ3010	AAATACCCCTGCTACCAGTCAGGCACAACAACTTATGCTGTTACTGAGACAACTTA
PEG2010 .	CCGACCGCCCCTGCAGTATTTCCATTGCTCAATACTTG-GCCACTTGTCTGGTGTTGAG
SEQ3001	A A A C C TH C TH C A C A C A C A C TH C C C C A A C TH C C C C A A C TH C C C C C A A C TH C C C C C A A C TH C C C C C C A A C TH C C C C C C C C C C C C C C C C C
SEQ3002	AAACCTGCTCAACACCAGACAAGTGGC-CAAGTATTGAGCAATGGAAATACTGCAGGGG
SEQ3003	AGACCTGCTCAACACCAGACGAGTGGC-CAAGTATTGAGTAATGGAAATACTGCAGGGG
SEQ3004	AGACCTGCTCAACACCAGACGAGTGGC-CAAGTATTGAGTAATGGAAATACTGCAGGGG
	AGACCTGCTCAACACCAGACGAGTGGC-CAAGTATTGAGTAATGGAAATACTGCAGGGG
SEQ3005	AGACCTGCTCAACACCAGACGAGTGGC-CAAGTATTGAGTAATGGAAATACTGCAGGGG
SEQ3007	AAACCTGCTCAACACCAGACAAGTGGC-CAAGTATTGAGCAATGGAAATACTGCAGGGG
SEQ3008	AGACCTGCTCAACACCAGACAAGTGGC-CAAGTATTGAGTAATGGAAATACTGCAGGGG
SEQ3009	AAACCTGCTCAACACCAGACAAGTGGC-CAAGTATTGAGCAATGGAAATACTGCAGGGG
SEQ3010	AGGTTTGTAAGTTGTCTCAGTAACAGCATAAGTTTGTTGTGCCTGACTGGTAGCAGGGG
SEQ3001	GGTCGGATCTGCTGCAGCACAAATGGCTGCTGCAACAGGAGTCCCTCAGTCTACTT
SEQ3002	TATTGGCTCAGCAGCTGCAGCACAAATGGCTGCCAACAGGAGTCCCTCAGTCTACTT
SEQ3003	TATTGGCTCAGCAGCAGCACAAATGGCTGCTGCAACAGGAGTCCCTCAGTCTACTT
SEQ3004	TATTGGCTCAGCAGCAGCACAAATGGCTGCTGCAACAGGAGTCCCTCAGTCTACTT
SEQ3005	TATTGGCTCAGCAGCTGCAGCACAAATGGCTGCTGCAACAGGAGTCCCTCAGTCTACTT
SEQ3007	GGTCGGATCTGCTGCAGCACAAATGGCTGCTGCAACAGGAGTCCCTCAGTCTACTT
SEQ3008	TATTGGCTCAGCAGCTGCAGCACAAATGGCTGCTGCAACAGGAGTCCCTCAGTCTACTT
SEQ3009	GGTCGGATCTGCTGCAGCACAAATGGCTGCTGCAACAGGAGTCCCTCAGTCTACTT
SEQ3010	A-TTTTCTGTTACAACTGCTTGTTCTACAGCCGCCTCTTCACTCGCAGTAACTTGTT
	The state of the s
SEQ3001	GGGAACATATTATTGCCCGTGAATCAAATGGTAATCCTAATGTTGCTAATGCCTCAGGAG
SEQ3002	GGGAACATATTATTGCCCGTGAATCAAATGGTAATCCTAATGTTGCTAATGCCTCAGGAG
SEQ3003 .	GGGAACATATTATTGCCCGTGAATCAAATGGTAATCCTAATGTTGCTAATGCCTCAGGAG
SEQ3004	GGGAACATATTATTGCCCGTGAATCAAATGGTAATCCTAATGTTGCTAATGCCTCAGGAG
SEQ3005	GGGAACATATTATTGCCCGTGAATCAAATGGTAATCCTAATGTTGCTAATGCCTCAGGAG
SEQ3007	GGGAACATATTATTGCCCGTGAATCAAATGGTAATCCTAATGTTGCTAATGCCTCAGGAG
SEQ3008	GGGAACATATTATTGCCCGTGAATCAAATGGTAATCCTAATGTTGCTAATGCCTCAGGAG
SEQ3009	CCCA CAMPATTA INCCCCCICA AT CAMAT GGTAAT CCTAAT GCTCAGGAG
SEQ3010	GGGAACATATTATTGCCCGTGAATCAAATGGTAATCCTAATGTTGCTAATGCCTCAGGAG
0110010	GCTGAGA-ATTAGCTTCTGTAGATTGAGAACTTGATTTTGGGGGCTTCATTAGATG
SEQ3001	CTTCAGGACTTTTCCAAACGATGCCAGGTTGGGGTTCAACAGCTACAGTTCAGGATCAAG
SEQ3002	CTTCAGGACTTTTCCAAACGATGCCAGGTTGGGGTTCAACAGCTACAGTTCAGGATCAAG
SEQ3003	CTTCAGGACTTTTCCAAACGATGCCAGGTTGGGGTTCAACAGCTACAGTTCAGGATCAAG
SEQ3004	CTTCAGGACTITICCAAACGAIGCCAGGITGGGGTTCAACAGCTACAGTTCAGGATCAAG CTTCAGGACTTTTCCAAACGATGCCAGGTTGGGGTTCAACAGCTACAGTTCAGGA
SEQ3005	CTTCAGGACTTTCCAAACGATGCCAGGTTGGGGTTCAACAGCTACAGTTCAGGA CTTCAGGACTTTTCCAAACGATGCCAGGTTGGGGTTCAACAGCTACAGTTCAGGATCAAG
SEQ3007	CTTCACACACTTICCAAACGATGCCAGGTTCAGGGTTCAACAGCTACAGTTCAGGATCAAG
SEQ3008	CTTCAGGACTTTTCCAAACGATGCCAGGTTGGGGTTCAACAGCTACAGTTCAGGATCAAG
SEQ3009	CTTCAGGACTTTTCCAAACGATGCCAGGTTGGGGTTCAACAGCTACAGTTCAGGATCAAG
	CTTCAGGACTTTTCCAAACGATGCCAGGTTGGGGTTCAACAGCTACAGTTCAGGATCAAG
SEQ3010	CCTGAGAAGGTTTTGGAGCCTGTTTTACATCTTCTACTTTTGATTTAGATGTCGC
SEQ3001	TA ATTICACCTATION A A COMMANDO COMO A COMMANDO A COMANDO A COMMANDO A COMMAND
\$EQ3001 SEQ3002	TAATTCAGCTATTAAAGCTTATCGTGCTCAAGGTTTATCAGCTTGGGGTTAC
SEQ3002	TAATTCAGCTATTAAAGCTTATCGTGCTCAAGGTTTATCAGCTTGGGGTTAC
	TAATTCAGCTATTAAAGCTTATCGTGCTCAAGGTTTATCAGCTTGGGGTTAC
SEQ3004	
SEQ3005	TAATTCAGCTATTAAAGCTTATCGTGCTCAAGGTTTATCAGCTTGGGGTTAC
SEQ3007	TAATTCAGCTATTAAAGCTTATCGTGCTCAAGGTTTATCAGCTTGGGGTTAC
SEQ3008	TAATTCAGCTATTAAAGCTT
SEQ3009	TAATTCAGCTATTAAAGCTTATCGTGCTCAAGGTTTATCAGCTTGGGGTTA
SEQ3010	TTAGTCA-TTTTTGATTTTTTGGCTACGCGAACTTTATCTGCTTTTGACAAAGA



### >SEQ ID NO 3050: 25\_1169NT frame: 1

KSSQVTTESLSKADKVRVAKKSKMTKATSKSKVEDVKQAPKPSQASNEVPKSSSQSTEAN SQQQVTASEEAAVEQAVVTENTPATSQAQQTYAVTETTYKPAQHQTSGQVLSNGNTAGAV GSAAAAQMAAATGVPQSTWEHIIARESNGNPNVANASGASGLFQTMPGWGSTATVQDQVN SAIKAYRAQGLSAWGY

#### >SEQ ID NO 3051:25\_18RS21 frame: 1

KSSQVTTESLSKADKVRVÄKKSKMTKATSKSKVEDVKQAPKPSQASNEAPKSSSQSTEAN SQQQVTASEEAAVEQAVVTENTPATSQAQQAYAVTETTYRPAQHQTSGQVLSNGNTAGAI GSAAAAQMAAATGVPQSTWEHIIARESNGNPNVANASGASGLFQTMPGWGSTATVQDQVN SAIKAYRAQGLSAWGY

### >SEQ ID NO 3052:25\_2603 frame: 1

KSSQVTTESLSKADKVRVÄKKSKMTKATSKSKVEDVKQAPKPSQASNEAPKSSSQSTEAN SQQQVTASEEAAVEQAVVTENTPATSQAQQAYAVTETTYRPAQHQTSGQVLSNGNTAGAI GSAAAAQMAAATGVPQSTWEHIIARESNGNPNVANASGASGLFQTMPGWGSTATVQDQVN SAIKAYRAQGLSAWGY

### >SEQ ID NO 3053:25\_090 frame: 3

AKKSKMIKATSKSKVEDVKQAPKPSQASNEAPKSSSQSTEANSQQQVTASEEAAVEQAVV TENTPATSQAQQAYAVTETTYRPAQHQTSGQVLSNGNTAGAIGSAAAAQMAAATGVPQST WEHIIARESNGNPNVANASGASGLFQTMPGWGSTATVQ

#### >SEQ ID NO 3054:25 A909 frame: 1

KATSKSKVEDVKQAPKPSQASNEAPKSSSQSTEANSQQQVTASEEAAVEQAVVTENTPAT SQAQQAYAVTETTYRPAQHQTSGQVLSNGNTAGAIGSAAAAQMAAATGVPQSTWEHIIAR ESNGNPNVANASGASGLFQTMPGWGSTATVQNQVNSAIKAYRAQGLS

#### >SEQ ID NO 3055:25 CJB110 frame: 3

SLSKADKVRVAKKSKMTKATSKSKVEDVKQAPKPSQASNEAPKSSSQSTEANSQQQVTAS EEAAVEQAVVTENTPATSQAQQAYAVTETTYRPAQHQTSGQVLSNGNTAGAIGSAAAAQM AAATGVPQSTWEHIIARESNGNPNVANASGASGLFQTMPGWGSTATVQDQVNSAIKAYRA QGLSAWGY

#### >SEQ ID NO 3056:25 COH1 frame: 1

KSSQVTTESLSKADKVRVÄKKSKMTKATSKSKVEDVKQAPKPSQASNEAPKSSSQSTEAN SQQQVTASEEAAVEQAVVTENTPATSQAQQTYAVTETTYKPAQHQTSGQVLSNGNTAGAV GSAAAAQMAAATGVPQSTWEHIIARESNGNPNVANASGASGLFQTMPGWGSTATVQDQVN SAIKAYRAQGLSAWGY

### >SEQ ID NO 3057:25\_H36B frame: 1

KSSQVTTESLSKADKVRVÄKKSKMTKATSKSKVEDVKQAPKPSQASNEAPKSSSQSTEAN SQQQVTASEEAAVEQAVVTENTPATSQAQQAYAVTETTYRPAQHQTSGQVLSNGNTAGAI GSAAAAQMAAATGVPQSTWEHIIARESNGNPNVANASGASGLFQTMPGWGSTATVQDQVN SAIKA

### >SEQ ID NO 3058:25\_M732 frame: 1

KSSQVTTESLSKADKVRVÄKKSKMTKATSKSKVEDVKQAPKPSQASNEAPKSSSQSTEAN SQQQVTASEEAAVEQAVVTENTPATSQAQQTYAVTETTYKPAQHQTSGQVLSNGNTAGAV GSAAAAQMAAATGVPQSTWEHIIARESNGNPNVANASGASGLFQTMPGWGSTATVQDQVN SAIKAYRAQGLSAWG

#### >SEQ ID NO 3059:25 M781 frame: 4

SLSKADKVRVAKKSKMTKĀTSKSKVEDVKQAPKPSQASNEAPKSSSQSTEANSQQQVTAS EEAAVEQAVVTENTPATSQAQQTYAVTETTYKPAQHQTSGQVLSNGNTAGAVGSAAAAQM AAATGVPQSTWEHIIARESNGNPNVANASGASGLFQTMPGWGSTATVQDQVNSAIKAYRA QGLSAWGY

SEQ3050	SSQVTTESLSKADKVRVAKKSKMTKATSKSKVEDVKQAPKPSQASNEVPKSSSQSTEAN
SEQ3051	SSQVTTESLSKADKVRVAKKSKMTKATSKSKVEDVKQAPKPSQASNEAPKSSSQSTEAN
SEQ3052	SSQVTTESLSKADKVRVAKKSKMTKATSKSKVEDVKQAPKPSQASNEAPKSSSQSTEAN
SEQ3053	AKKSKMIKATSKSKVEDVKQAPKPSQASNEAPKSSSQSTEAN
SEQ3054	Katskskvedvkqapkpsqasneapksssqstean
SEQ3055	SLSKADKVRVAKKSKMTKATSKSKVEDVKQAPKPSQASNEAPKSSSQSTEAN
SEQ3056	SSQVTTESLSKADKVRVAKKSKMTKATSKSKVEDVKQAPKPSQASNEAPKSSSQSTEAN
SEQ3057	SSQVTTESLSKADKVRVAKKSKMTKATSKSKVEDVKQAPKPSQASNEAPKSSSQSTEAN
SEQ3058	SSQVTTESLSKADKVRVAKKSKMTKATSKSKVEDVKQAPKPSQASNEAPKSSSQSTEAN
SEQ3059	SLSKADKVRVAKKSKMTKATSKSKVEDVKOAPKPSOASNEAPKSSSOSTEAN

604062 1082602

# Table 30: Comparative Sequences relating to SA62.47 (protein of uknown function / lip protein, putative)

SEQ3050	SQQQVTASEEAAVEQAVVTENTPATSQAQQTYAVTETTYKPAQHQTSGQVLSNGNTAGAV
SEQ3051	SQQQVTASEEAAVEQAVVTENTPATSQAQQAYAVTETTYRPAQHQTSGQVLSNGNTAGAI
SEQ3052	SQQQVTASEEAAVEQAVVTENTPATSQAQQAYAVTETTYRPAQHQTSGQVLSNGNTAGAI
SEQ3053	SQQQVTASEEAAVEQAVVTENTPATSQAQQAYAVTETTYRPAQHQTSGQVLSNGNTAGAI
SEQ3054	SQQQVTASEEAAVEQAVVTENTPATSQAQQAYAVTETTYRPAQHQTSGQVLSNGNTAGAI
SEQ3055	SQQQVTASEEAAVEQAVVTENTPATSQAQQAYAVTETTYRPAQHQTSGQVLSNGNTAGAI
SEQ3056	SQQQVTASEEAAVEQAVVTENTPATSQAQQTYAVTETTYKPAQHQTSGQVLSNGNTAGAV
SEQ3057	SQQQVTASEEAAVEQAVVTENTPATSQAQQAYAVTETTYRPAQHQTSGQVLSNGNTAGAI
SEQ3058	SQQQVTASEEAAVEQAVVTENTPATSQAQQTYAVTETTYKPAQHQTSGQVLSNGNTAGAV
SEQ3059	SQQQVTASEEAAVEQAVVTENTPATSQAQQTYAVTETTYKPAQHQTSGQVLSNGNTAGAV
SEQ3050	GSAAAAQMAAATGVPQSTWEHIIARESNGNPNVANASGASGLFQTMPGWGSTATVQDQVN
SEQ3051	GSAAAAQMAAATGVPQSTWEHIIARESNGNPNVANASGASGLFQTMPGWGSTATVQDQVN
SEQ3052	GSAAAAQMAAATGVPQSTWEHIIARESNGNPNVANASGASGLFQTMPGWGSTATVQDQVN
SEQ3053	GSAAAAQMAAATGVPQSTWEHIIARESNGNPNVANASGASGLFQTMPGWGSTATVQ
SEQ3054	GSAAAAQMAAATGVPQSTWEHIIARESNGNPNVANASGASGLFQTMPGWGSTATVQNQVN
SEQ3055	GSAAAAQMAAATGVPQSTWEHIIARESNGNPNVANASGASGLFQTMPGWGSTATVQDQVN
SEQ3056	GSAAAAQMAAATGVPQSTWEHIIARESNGNPNVANASGASGLFQTMPGWGSTATVQDQVN
SEQ3057	GSAAAAQMAAATGVPQSTWEHIIARESNGNPNVANASGASGLFQTMPGWGSTATVQDQVN
SEQ3058	GSAAAAQMAAATGVPQSTWEHIIARESNGNPNVANASGASGLFQTMPGWGSTATVQDQVN
SEQ3059	GSAAAAQMAAATGVPQSTWEHIIARESNGNPNVANASGASGLFQTMPGWGSTATVQDQVN
SEQ3050	AIKAYRAQGLSAWGY
SEQ3051	AIKAYRAQGLSAWGY
SEQ3052	AIKAYRAQGLSAWGY
SEQ3053	
SEQ3054	AIKAYRAQGLS
SEQ3055	AIKAYRAQGLSAWGY
SEQ3056	AIKAYRAQGLSAWGY
SEQ3057	AIKA
SEQ3058	AIKAYRAQGLSAWG-
SEQ3059	AIKAYRAQGLSAWGY



# Table 51: C mparative Sequences relating to SA 52148 (LysM domain protein)

SEQ ID NO. 3101: SAG2148 FROM THE 1169NT1 GBS TYPE V STRAIN
GCATCTTATACCGTGAAATCAGGTGATACCTTATCAGCTATTGCTAAAAATCATAAAACTACGGTACAAGAGTTAGTG
TCTCTCAATAGTATCAGTAACGCTGATGTCATCAGTATAGGTGATGTTTTAAAATTGGATAATTCTACAGCTAGTCAA
GCAGAAGCAAAATCTCAACCAACAATTGAAAATTCAATGAATTCTTCATCAAATTTGAGTTCAAGTGATTCAGCTGCA
AAAGAAGAAAATAGCTCGTCGTGAATCAAATGGTAGTTATACTGCACAGAATGGACAATATTATGGAAGATATCAACTG
TCTCAATCTTACCTAAATGGCGACTTATCTCCTGAAAATCAAGAAAAAGTAGCGGACAATTATGTGGCTTCTCGTTAC
GGATCTTGGTCGGCAGCGCTATCATTTTGGAATAGTAACGGCTGGTAT

SEQ ID NO. 3102: SAG2148 FROM THE 18RS21 GBS TYPE II STRAIN
GCATCTTATACCGTGAAATCAGGTGATACCTTATCAGCTATTGCTAAAAATCATAAAACTACGGTACAAGAGTTAGTG
TCTCTCAATAGTATCAGTAACGCTGATGTCATCAGTATAGGTGATGTTTTAAAATTGGATAATTCTACAGCTAGTCAA
GCAGAAGCAAAATCTCAACCAACAATTGAAAATTCAATGAATTCTACAGAATTTGAGTTCAAGTGATTCAGCCGCA
AAAGAAGAAAATAGCTCGTCGTGAATCAAATGGTAGTTATACTGCACAGAATGGACAATATTATGGAAGATATCAACTG
TCTCAATCTTACCTAAATGGCGACTTATCTCCTGAAAATCAAGAAAAAGTAGCGGACAATTATGTGGTTTCTCGTTAC
GGATCTTGGTCGGCAGCGCTATCATTTTGGAATAGTAACGGCTGGTAT

SEQ ID NO. 3103: SAG2148 FROM THE 2603 V/R GBS TYPE V STRAIN
GCATCTTATACCGTGAAATCAGGTGATACCTTATCAGCTATTGCTAAAAATCATAAAACTACGGTACAAGAGTTAGTG
TCTCTCAATAGTATCAGTAACGCTGATGTCATCAGTATAGGTGATGTTTTAAAATTGGATAATTCTACAGCTAGTCAA
GCAGAAGCAAAATCTCAACCAACAATTGAAAATTCAATGAATTCTTCATCAAATTTGAGTTCAAGTGATTCAGCCGCA
AAAGAAGAAAATAGCTCGTCGTGAATCAAATGGTAGTTATACTGCACAGAATGGACAATTATTGGAAGATATCAACTG
TCTCAATCTTACCTAAATGGCGACTTATCTCCTGAAAATCAAGAAAAAGTAGCGGACAATTATGTGGTTTCTCGTTAC
.GGATCTTGGTCGGCAGCGCTATCATTTTGGAATAGTAACGGCTGGTAT

SEQ ID NO. 3104: SAG2148 FROM THE 090 GBS TYPE IA STRAIN
GCATCTTATACCGTGAAATCAGGTGATACCTTATCAGCTATTGCTAAAAATCATAAAACTACGGTACAAGAGTTAGTG
TCTCTCAATAGTATCAGTAACGCTGATGTCATCAGTATAGGTGATGTTTTAAAATTTGGATAATTCTAAAGCTAGTCAA
GCAGAAGCAAAATCTCAACCAACAATTGAAAATTCAATGAATTCTTCATCAAATTTGAGTTCAAGTGATTCAGCCGCA
AAAGAAGAAAATAGCTCGTCGTGAATCAAATGGTAGTTATACTGCACAGAATGGACAATATTATGGAAGATATCAACTG
TCTCAATCTTACCTAAATGGCGACTTATCTCCTGAAAATCAAGAAAAAGTAGCGGACAATTATGTGGTTTCTCGTTAC
GGATCTTGGTCGGCAGCGCTATCATTTTGGAATAGTAACGGCTGGTAT

SEQ ID NO. 3105: SAG2148 FROM THE A909 GBS TYPE Ia STRAIN
GCATCTTATACCGTGAAATCAGGTGATACCTTATCAGCTATTGCTAAAAATCATAAAACTACGGTACAAGAGTTAGTG
TCTCTCAATAGTATCAGTAACGCTGATGTCATCAGTATAGGTGATGTTTTTAAAATTGGATAATTCTACAGCTAGTCAA
GCAGAAGCAAAATCTCAACCAACAATTGAAAATTCAATGAATTCTTCATCAAATTTGAGTTCAAGTGATTCAGCCGCA
AAAGAAGAAAATAGCTCGTCGTGAATCAAATGGTAGTTATACTGCACAGAATGGACAATATTATGGAAGATATCAACTG
TCTCAATCTTACCTAAATGGCGACTTATCTCCTGAAAATCAAGAAAAAGTAGCGGACAATTATGTGGCTTCTCGTTAC
GGATCTTGGTCGGCAGCGCTATCATTTTGGAATAGTAACGGCTGGTAT

SEQ ID NO. 3106: SAG2148 FROM THE CJB110 GBS NONTYPEABLE STRAIN
GCATCTTATACCGTGAAATCAGGTGATACCTTATCAGCTATTGCTAAAAATCATAAAACTACGGTACAAGAGTTAGTG
TCTCTCAATAGTATCAGTAACGCTGATGTCATCAGTATAGGTGATGTTTTAAAATTGGATAATTCTAAAGCTAGTCAA
GCAGAAGCAAAATCTCAACCAACAATTGAAAATTCAATGAATTCTACACAAATTTGAGTTCAAGTGATTCAGCCGCA
AAAGAAGAAAATAGCTCGTCGTGAATCAAATGGTAGTTATACTGCACAGAATGGACAATTTATGGAAGATATCAACTG
TCTCAATCTTACCTAAATGGCGACTTATCTCCTGAAAATCAAGAAAAAGTAGCGGACAATTATGTGGTTTCTCGTTAC
GGATCTTGGTCGGCAGCGCTATCATTTTGGAATAGTAACGGCTGGTAT

# Table 31: Comparative Sequences relating to SA 52148 (LysM d main pr tein)

# SEQ ID NO. 3108: SAG2148 FROM THE H36b GBS TYPE ID STRAIN (REVERSE COMPLEMENT)

GCATCTTATACCGTGAAATCAGGTGATACCTTATCAGCTATTGCTAAAAATCATAAAACTACGGTACAAGAGTTAGTG
TCTCTCAATAGTATCAGTAACGCTGATGTCATCAGTATAGGTGATGTTTTAAAATTGGATAATTCTACAGCTAGTCAA
GCAGAAGCAAAATCTCAACCAACAATTGAAAATTCAATGAATTCTTCATCAAATTTGAGTTCAAGTGATTCAGCCGCA
AAAGAAGAAAATAGCTCGTCGTGAAATCAAATGGTAGTTATACTGCACAGAATGGACAATTATTATGGAAGATATCAACTG
TCTCAATCTTACCTAAATGGCGACTTATCTCCTGAAAATCAAGAAAAAGTAGCGGACAATTATGTGGCTTCTCGTTAC
GGATCTTGGTCGGCAGCGCTATCATTTTGGAATAGTAACGGCTGGTAT

# SEQ ID NO. 3109: SAG2148 FROM THE JM9130013 GBS TYPE VIII STRAIN (REVERSE COMPLEMENT)

GCATCTTATACCGTGAAATCAGGTGATACCTTATCAGCTATTGCTAAAAATCATAAAACTACGGTACAAGAGTTAGTG
TCTCTCAATAGTATCAGTAACGCTGACGTCATCAGTATAGGTGATGTTTTTAAAATTGGATAATTCTACAACTAGTCAA
GCAGAAGCAAAATCTCAACCAACAATTGAAAATTCAATGAATTCTTCATCAAATTTGAGTTCAAGTGATTCAGCCGCA
AAAGAAGAAATAGCTCGTCGTGAATCAAATGGTAGTTATACTGCACAGAATGGACAATATTATGGAAGATATCAACTG
TCTCAATCTTACCTAAATGGCGACTTATCTCCTGAAAATCAAGAAAAAGTAGCGGACAATTATGTGGCTTCTCGTTAC
GGATCTTGGTCGGCAGCGCTATCATTTTGGAATAGTAACGGCTGGTAT

#### SEQ ID NO. 3110: SAG2148 FROM THE M732 GBS TYPE III STRAIN

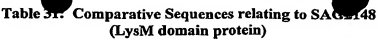
# SEQ ID NO. 3111: SAG2148 FROM THE M781 GBS TYPE III STRAIN (REVERSE COMPLEMENT)

SEQ3101	GCATCTTATACCGTGAAATCAGGTGATACCTTATCAGCTATTGCTAAAAATCATAAAACT
SEQ3102	GCATCTTATACCGTGAAATCAGGTGATACCTTATCAGCTATTGCTAAAAATCATAAAACT
SEQ3103	GCATCTTATACCGTGAAATCAGGTGATACCTTATCAGCTATTGCTAAAAATCATAAAACT
SEQ3104	GCATCTTATACCGTGAAATCAGGTGATACCTTATCAGCTATTGCTAAAAATCATAAAACT
SEQ3105	GCATCTTATACCGTGAAATCAGGTGATACCTTATCAGCTATTGCTAAAAATCATAAAACT
SEQ3106	GCATCTTATACCGTGAAATCAGGTGATACCTTATCAGCTATTGCTAAAAATCATAAAACT
SEQ3107	GCATCTTATACCGTGAAATCAGGTGATACCTTATCAGCTATTGCTAAAAATCATAAAACT
SEQ3108	GCATCTTATACCGTGAAATCAGGTGATACCTTATCAGCTATTGCTAAAAATCATAAAACT
SEQ3109	GCATCTTATACCGTGAAATCAGGTGATACCTTATCAGCTATTGCTAAAAATCATAAAACT
SEQ3110	GCATCTTATACCGTGAAATCAGGTGATACCTTATCAGCTATTGCTAAAAATCATAAAACT
SEQ3111	GCATCTTATACCGTGAAATCAGGTGATACCTTATCAGCTATTGCTAAAAATCATAAAACT
SEQ3101	ACGGTACAAGAGTTAGTGTCTCTCAATAGTATCAGTAACGCTGATGTCATCAGTATAGGT
SEQ3102	ACGGTACAAGAGTTAGTGTCTCTCAATAGTATCAGTAACGCTGATGTCATCAGTATAGGT
SEQ3103	ACGGTACAAGAGTTAGTGTCTCTCAATAGTATCAGTAACGCTGATGTCATCAGTATAGGT
SEQ3104	ACGGTACAAGAGTTAGTGTCTCTCAATAGTATCAGTAACGCTGATGTCATCAGTATAGGT
SEQ3105	ACGGTACAAGAGTTAGTGTCTCTCAATAGTATCAGTAACGCTGATGTCATCAGTATAGGT
SEQ3106	ACGGTACAAGAGTTAGTGTCTCTCAATAGTATCAGTAACGCTGATGTCATCAGTATAGGT
SEQ3107	ACGGTACAATAGTTAGTGTCTCTCAATAGTATCAGTAACGCTGATGTCATCAGTATAGGT
SEQ3108	ACGGTACAAGAGTTAGTGTCTCTCAATAGTATCAGTAACGCTGATGTCATCAGTATAGGT
SEQ3109	ACGGTACAAGAGTTAGTGTCTCTCAATAGTATCAGTAACGCTGACGTCATCAGTATAGGT
SEQ3110	ACGGTACAATAGTTAGTGTCTCTCAATAGTATCAGTAACGCTGATGTCATCAGTATAGGT
SEQ3111	ACGGTACAATAGTTAGTGTCTCTCAATAGTATCAGTAACGCTGATGTCATCAGTATAGGT



# Table 31: Comparative Sequences relating to SA 52148 (LysM domain protein)

SEQ3101	GATGTTTTAAAATTGGATAATTCTACAGCTAGTCAAGCAGAAGCAAAATCTCAACCAAC
SEQ3102	GATGTTTTAAAATTGGATAATTCTACAGCTAGTCAAGCAGAAGCAAAATCTCAACCAAC
SEQ3103	GATGTTTTAAAATTGGATAATTCTACAGCTAGTCAAGCAGAAGCAAAATCTCAACCAAC
SEQ3104	GATGTTTTAAAATTGGATAATTCTAAAGCTAGTCAAGCAGAAGCAAAATCTCAACCAAC
SEQ3105	GATGTTTTAAAATTGGATAATTCTACAGCTAGTCAAGCAGAAGCAAAATCTCAACCAAC
SEQ3106	GATGTTTTAAAATTGGATAATTCTAAAGCTAGTCAAGCAGAAGCAAAATCTCAACCAAC
SEQ3107	GATGTTTTAAAATTGGATAATTCTACAGCTAGTCAAGCAGAAGCAAAATCTCAACCAAC
SEQ3108	GATGTTTTAAAATTGGATAATTCTACAGCTAGTCAAGCAGAAGCAAAATCTCAACCAAC
SEQ3109	GATGTTTTAAAATTGGATAATTCTACAACTAGTCAAGCAGAAGCAAAATCTCAACCAAC
SEQ3110	GATGTTTTAAAATTGGATAATTCTACAGCTAGTCAAGCAGAAGCAAAATCTCAACCAAC
SEQ3111	GATGTTTTAAAATTGGATAATTCTACAGCTAGTCAAGCAGAAGCAAAATCTCAACCAAC
CPA21A1	AMMCA A A AMMCA AMCA AMMCA AMMCA AMACA A AMMCA AMMCA ACMACA ACA
SEQ3101 SEQ3102	ATTGAAAATTCAATGAATTCTTCATCAAATTTGAGTTCAAGTGATTCAGCTGCAAAAGAA ATTGAAAATTCAATGAATTCTTCATCAAATTTGAGTTCAAGTGATTCAGCCGCAAAAGAA
SEQ3103	ATTGAAAATTCAATGAATTCTTCATCAAATTTGAGTTCAAGTGATTCAGCCGCAAAAGAA ATTGAAAATTCAATGAATTCTTCATCAAATTTGAGTTCAAGTGATTCAGCCGCAAAAGAA
SEQ3104	ATTGAAAATTCAATGAATTCTTCATCAAATTTGAGTTCAAGTGATTCAGCCGCAAAAGAA ATTGAAAATTCAATGAATTCTTCATCAAATTTGAGTTCAAGTGATTCAGCCGCAAAAGAA
SEQ3105	ATTGAAAATTCAATGAATTCTTCATCAAATTTGAGTTCAAGTGATTCAGCCGCAAAAGAA ATTGAAAATTCAATGAATTCTTCATCAAAATTTGAGTTCAAGTGATTCAGCCGCAAAAGAA
SEQ3106	ATTGAAAATTCAATGAATTCTTCATCAAATTTGAGTTCAAGTGATTCAGCCGCAAAAGAA ATTGAAAATTCAATGAATTCTTCATCAAATTTGAGTTCAAGTGATTCAGCCGCAAAAGAA
SEQ3107	ATTGAAAATTCAATGAATTCTTCATCAAATTTGAGTTCAAGTGATTCAGCCGCAAAAGAA ATTGAAAATTCAATGAATTCTTCATCAAAATTTGAGTTCAAGTGATTCAGCTGCAAAAGAA
SEQ3108	ATTGAAAATTCAATGAATTCTTCATCAAATTTGAGTTCAAGTGATTCAGCCGCAAAAGAA
SEQ3109	ATTGAAAATTCAATGAATTCTTCATCAAATTTGAGTTCAAGTGATTCAGCCGCAAAAGAA
SEQ3110	ATTGAAAATTCAATGAATTCTTCATCAAATTTGAGTTCAAGTGATTCAGCCGCAAAAGAA
SEQ3111	ATTGAAAATTCAATGAATTCTTCATCAAATTTGAGTTCAAGTGATTCAGCTGCAAAAGAA
<b></b>	wasser wasser to to out out the transfer to the transfer
SEQ3101	GAAATAGCTCGTCGTGAATCAAATGGTAGTTATACTGCACAGAATGGACAATATTATGGA
SEQ3102	GAAATAGCTCGTCGTGAATCAAATGGTAGTTATACTGCACAGAATGGACAATATTATGGA
SEQ3103	GAAATAGCTCGTCGAATCAAATGGTAGTTATACTGCACAGAATGGACAATATTATGGA
SEQ3104	GAAATAGCTCGTCGTGAATCAAATGGTAGTTATACTGCACAGAATGGACAATATTATGGA
SEQ3105	GAAATAGCTCGTCGAATCAAATGGTAGTTATACTGCACAGAATGGACAATATTATGGA
SEQ3106	GAAATAGCTCGTCGTGAATCAAATGGTAGTTATACTGCACAGAATGGACAATATTATGGA
SEQ3107	GAAATAGCTCGTCGTGAATCAAATGGTAGTTATACTGCACAGAATGGACAATATTATGGA
SEQ3108	GAAATAGCTCGTCGTGAATCAAATGGTAGTTATACTGCACAGAATGGACAATATTATGGA
SEQ3109	GAAATAGCTCGTCGTGAATCAAATGGTAGTTATACTGCACAGAATGGACAATATTATGGA
SEQ3110	GAAATAGCTCGTCGTGAATCAAATGGTAGTTATACTGCACAGAATGGACAATATTATGGA
SEQ3111	GAAATAGCTCGTCGTGAATCAAATGGTAGTTATACTGCACAGAATGGACAATATTATGGA
SEQ3101	AGATATCAACTGTCTCAATCTTACCTAAATGGCGACTTATCTCCTGAAAATCAAGAAAAA
SEQ3102	AGATATCAACTGTCTCAATCTTACCTAAATGGCGACTTATCTCCTGAAAATCAAGAAAAA AGATATCAACTGTCTCAATCTTACCTAAATGGCGACTTATCTCCTGAAAATCAAGAAAAA
· SEQ3103	AGATATCAACTGTCTCAATCTTACCTAAATGGCGACTTATCTCCTGAAAATCAAGAAAAA AGATATCAACTGTCTCAATCTTACCTAAATGGCGACTTATCTCCTGAAAATCAAGAAAAA
SEQ3104	AGATATCAACTGTCTCAATCTTACCTAAATGGCGACTTATCTCCTGAAAATCAAGAAAAA
SEQ3105	AGATATCAACTGTCTCAATCTTACCTAAATGGCGACTTATCTCCTGAAAATCAAGAAAAA
SEQ3106	AGATATCAACTGTCTCAATCTTACCTAAATGGCGACTTATCTCCTGAAAATCAAGAAAAA
SEQ3107	AGATATCAACTGTCTCAATCTTACCTAAATGGCGACTTATCTCCTGAAAATCAAGAAAAA
SEQ3108	AGATATCAACTGTCTCAATCTTACCTAAATGGCGACTTATCTCCTGAAAATCAAGAAAA
SEQ3109	AGATATCAACTGTCTCAATCTTACCTAAATGGCGACTTATCTCCTGAAAATCAAGAAAAA
SEQ3110	AGATATCAACTGTCTCAATCTTACCTAAATGGCGACTTATCTCCTGAAAATCAAGAAAAA
SEQ3111	AGATATCAACTGTCTCAATCTTACCTAAATGGCGACTTATCTCCTGAAAATCAAGAAAAA
SEQ3101	GTAGCGGACAATTATGTGGCTTCTCGTTACGGATCTTGGTCGGCAGCGCTATCATTTTGG
SEQ3102	GTAGCGGACAATTATGTGGTTTCTCGTTACGGATCTTGGTCGGCAGCGCTATCATTTTGG
SEQ3103	GTAGCGGACAATTATGTGGTTTCTCGTTACGGATCTTGGTCGGCAGCGCTATCATTTTGG
SEQ3104	GTAGCGGACAATTATGTGGTTTCTCGTTACGGATCTTGGTCGGCAGCGCTATCATTTTGG
SEQ3105	GTAGCGGACAATTATGTGGCTTCTCGTTACGGATCTTGGTCGGCAGCGCTATCATTTTGG
SEQ3106	GTAGCGGACAATTATGTGGTTTCTCGTTACGGATCTTGGTCGGCAGCGCTATCATTTTGG
SEQ3107	GTAGCGGACAATTATGTGGCTTCTCGTTACGGATCTTGGTCGGCAGCGCTATCATTTTGG
SEQ3108	GTAGCGGACAATTATGTGGCTTCTCGTTACGGATCTTGGTCGGCAGCGCTATCATTTTGG
SEQ3109	GTAGCGGACAATTATGTGGCTTCTCGTTACGGATCTTGGTCGGCAGCGCTATCATTTTGG
SEQ3110	GTAGCGGACAATTATGTGGCTTCTCGTTACGGATCTTGGTCGGCAGCGCTATCATTTTGG
SEQ3111	GTAGCGGACAATTATGTGGCTTCTCGTTACGGATCTTGGTCGGCAGCGCTATCATTTTGG



SEQ3101	AATAGTAACGGCTGGTAT
SEQ3102	AATAGTAACGGCTGGTAT
SEQ3103	AATAGTAACGGCTGGTAT
SEQ3104	AATAGTAACGGCTGGTAT
SEQ3105	AATAGTAACGGCTGGTAT
SEQ3106	AATAGTAACGGCTGGTAT
SEQ3107	AATAGTAACGGCTGGTAT
SEQ3108	<b>AATAGTAACGGCTGGŤAT</b>
SEQ3109	AATAGTAACGGCTGGTAT
SEQ3110	AATAGTAACGGCTGGTAT
SEQ3111	AATAGTAACGGCTGGTAT

#### >SEQ ID NO 3150:15\_1169NT frame: 1

ASYTVKSGDTLSAIAKNHKTTVQELVSLNSISNADVISIGDVLKLDNSTASQAEAKSQPT IENSMNSSSNLSSSDSAAKEEIARRESNGSYTAQNGQYYGRYQLSQSYLNGDLSPENQEK VADNYVASRYGSWSAALSFWNSNGWY

#### >SEQ ID NO 3151:15\_18R\$21 frame: 1

ASYTVKSGDTLSAIAKNHKTTVQELVSLNSISNADVISIGDVLKLDNSTASQAEAKSQPT IENSMNSSSNLSSSDSAAKEEIARRESNGSYTAQNGQYYGRYQLSQSYLNGDLSPENQEK· VADNYVVSRYGSWSAALSFWNSNGWY

#### >SEQ ID NO 3152:15 2603 frame: 1

ASYTVKSGDTLSAIAKNHKTTVQELVSLNSISNADVISIGDVLKLDNSTASQAEAKSQPT IENSMNSSSNLSSSDSAAKEEIARRESNGSYTAQNGQYYGRYQLSQSYLNGDLSPENQEK VADNYVVSRYGSWSAALSFWNSNGWY

#### >SEQ ID NO 3153:15 090 frame: 1

ASYTVKSGDTLSATAKNHKTTVQELVSLNSISNADVISIGDVLKLDNSKASQAEAKSQPT IENSMNSSSNLSSSDSAAKEEIARRESNGSYTAQNGQYYGRYQLSQSYLNGDLSPENQEK VADNYVVSRYGSWSAALSFWNSNGWY

#### >SEQ ID NO 3154:15\_A909 frame: 1

ASYTVKSGDTLSAIAKNHKTTVQELVSLNSISNADVISIGDVLKLDNSTASQAEAKSQPT IENSMNSSSNLSSSDSAAKEEIARRESNGSYTAQNGQYYGRYQLSQSYLNGDLSPENQEK VADNYVASRYGSWSAALSFWNSNGWY

#### >SEQ ID NO 3155:15 CJB110 frame: 1

ASYTVKSGDTLSAIAKNHKTTVQELVSLNSISNADVISIGDVLKLDNSKASQAEAKSQPT IENSMNSSSNLSSSDSAAKEEIARRESNGSYTAQNGQYYGRYQLSQSYLNGDLSPENQEK VADNYVVSRYGSWSAALSFWNSNGWY

#### >SEQ ID NO 3156:15\_COH1 frame: 1

ASYTVKSGDTLSAIAKNHKTTVQ.LVSLNSISNADVISIGDVLKLDNSTASQAEAKSQPT IENSMNSSSNLSSSDSAAKEEIARRESNGSYTAQNGQYYGRYQLSQSYLNGDLSPENQEK VADNYVASRYGSWSAALSFWNSNGWY

#### >SEQ ID NO 3157:15\_H36B frame: 1

ASYTVKSGDTLSAIAKNHKTTVQELVSLNSISNADVISIGDVLKLDNSTASQAEAKSQPT IENSMNSSSNLSSSDSAAKEEIARRESNGSYTAQNGQYYGRYQLSQSYLNGDLSPENQEK VADNYVASRYGSWSAALSFWNSNGWY

#### >SEQ ID NO 3158:15\_JM9130013 frame: 1

ASYTVKSGDTLSAIAKNHKTTVQELVSLNSISNADVISIGDVLKLDNSTTSQAEAKSQPT IENSMNSSSNLSSSDSAAKEEIARRESNGSYTAQNGQYYGRYQLSQSYLNGDLSPENQEK VADNYVASRYGSWSAALSFWNSNGWY

#### >SEQ ID NO 3159:15\_M732 frame: 1

ASYTVKSGDTLSAIAKNHKTTVQ.LVSLNSISNADVISIGDVLKLDNSTASQAEAKSQPT IENSMNSSSNLSSSDSAAKEEIARRESNGSYTAQNGQYYGRYQLSQSYLNGDLSPENQEK VADNYVASRYGSWSAALSFWNSNGWY

#### >SEQ ID NO 3160:15 M781 frame: 1

ASYTVKSGDTLSAIAKNHKTTVQ.LVSLNSISNADVISIGDVLKLDNSTASQAEAKSQPT IENSMNSSSNLSSSDSAAKEEIARRESNGSYTAQNGQYYGRYQLSQSYLNGDLSPENQEK VADNYVASRYGSWSAALSFWNSNGWY





# Table 31: Comparative Sequences relating to SAG2148 (LysM domain protein)

SEQ3150	ASYTVKSGDTLSAIAKNHKTTVQELVSLNSISNADVISIGDVLKLDNSTASQAEAKSQPT
SEQ3151	ASYTVKSGDTLSAIAKNHKTTVQELVSLNSISNADVISIGDVLKLDNSTASQAEAKSQPT
SEQ3152	ASYTVKSGDTLSAIAKNHKTTVQELVSLNSISNADVISIGDVLKLDNSTASQAEAKSQPT
SEQ3153	ASYTVKSGDTLSAIAKNHKTTVQELVSLNSISNADVISIGDVLKLDNSKASQAEAKSQPT
SEQ3154	ASYTVKSGDTLSAIAKNHKTTVQELVSLNSISNADVISIGDVLKLDNSTASQAEAKSQPT
SEQ3155	ASYTVKSGDTLSAIAKNHKTTVQELVSLNSISNADVISIGDVLKLDNSKASQAEAKSQPT
SEQ3156	ASYTVKSGDTLSAIAKNHKTTVQ-LVSLNSISNADVISIGDVLKLDNSTASQAEAKSQPT
SEQ3157	ASYTVKSGDTLSAIAKNHKTTVQELVSLNSISNADVISIGDVLKLDNSTASQAEAKSQPT
SEQ3158	ASYTVKSGDTLSAIAKNHKTTVQELVSLNSISNADVISIGDVLKLDNSTTSQAEAKSQPT
SEQ3159	ASYTVKSGDTLSAIAKNHKTTVQ-LVSLNSISNADVISIGDVLKLDNSTASQAEAKSQPT
SEQ3160	ASYTVKSGDTLSAIAKNHKTTVQ-LVSLNSISNADVISIGDVLKLDNSTASQAEAKSQPT
SEQ3150	IENSMNSSSNLSSSDSAAKEEIARRESNGSYTAQNGQYYGRYQLSQSYLNGDLSPENQEK
SEQ3151	IENSMNSSSNLSSSDSAAKEEIARRESNGSYTAQNGQYYGRYQLSQSYLNGDLSPENQEK
SEQ3152	IENSMNSSSNLSSSDSAAKEEIARRESNGSYTAQNGQYYGRYQLSQSYLNGDLSPENQEK
SEQ3153	IENSMNSSSNLSSSDSAAKEEIARRESNGSYTAQNGQYYGRYQLSQSYLNGDLSPENQEK
SEQ3154	IENSMNSSSNLSSSDSAAKEEIARRESNGSYTAQNGQYYGRYQLSQSYLNGDLSPENQEK
SEQ3155	IENSMNSSSNLSSSDSAAKEEIARRESNGSYTAQNGQYYGRYQLSQSYLNGDLSPENQEK
SEQ3156	IENSMNSSSNLSSSDSAAKEEIARRESNGSYTAQNGQYYGRYQLSQSYLNGDLSPENQEK
SEQ3157	IENSMNSSSNLSSSDSAAKEEIARRESNGSYTAQNGQYYGRYQLSQSYLNGDLSPENQEK
SEQ3158	IENSMNSSSNLSSSDSAAKEEIARRESNGSYTAQNGQYYGRYQLSQSYLNGDLSPENQEK
SEQ3159	IENSMNSSSNLSSSDSAAKEEIARRESNGSYTAQNGQYYGRYQLSQSYLNGDLSPENQEK
SEQ3160	IENSMNSSSNLSSSDSAAKEEIARRESNGSYTAQNGQYYGRYQLSQSYLNGDLSPENQEK
SEQ3150	VADNYVASRYGSWSAALSFWNSNGWY
SEQ3151	VADNYVVSRYGSWSAALSFWNSNGWY
SEQ3152	VADNYVVSRYGSWSAALSFWNSNGWY
SEQ3153	VADNYVVSRYGSWSAALSFWNSNGWY
SEQ3154	VADNYVASRYGSWSAALSFWNSNGWY
SEQ3155	VADNYVVSRYGSWSAALSFWNSNGWY
SEQ3156	VADNYVASRYGSWSAALSFWNSNGWY
SEQ3157	VADNYVASRYGSWSAALSFWNSNGWY
SEQ3158	VADNYVASRYGSWSAALSFWNSNGWY
SEQ3159	VADNYVASRYGSWSAALSFWNSNGWY
SEQ3160	VADNYVASRYGSWSAALSFWNSNGWY



Tabl 32: Conv rsion of ORF R f N s. with SAG Ref Nos.

ORF Ref N .	SAGxxxx Ref No.	aa	Annotation
ORF00003	SAG0017	447	PcsB protein
ORF00004	SAG0018	322	ribose-phosphate pyrophosphokinase
ORF00005	SAG0019	391	aminotransferase, class I
ORF00006	SAG0020	253	recombination protein O
ORF00008	SAG0021	283	protease, putative
ORF00009	SAG0022	330	fatty acid/phospholipid synthesis protein PlsX
ORF00010	SAG0023	79	acyl carrier protein
ORF00011	SAG0024	234	phosphoribosylaminoimidazole-succinocarboxamide
0.4.00017	0/100024	254	synthase
ORF00012	SAG0025	1241	phosphoribosylformylglycinamidine synthase, putative
ORF00013	SAG0026	484	amidophosphoribosyltransferase
ORF00014	SAG0027	340	phosphoribosylformylglycinamidine cyclo-ligase
ORF00015	SAG0028	182	phosphoribosylglycinamide formyltransferase
ORF00016	SAG0029	250	acetyltransferase, GNAT family
ORF00017	SAG0030	515	phosphoribosylaminoimidazolecarboxamide
0141 00017	OACOUSO	3.3	formyltransferase/IMP cyclohydrolase
ORF00018	SAG0031	283	peptidase, M23/M37 family
ORF00020	SAG0032	434	group B streptococcal surface immunogenic protein
			group a choptococci curiuso immenoganio protein
ORF00021	SAG0033	232	N-acetylmannosamine-6-P epimerase, putative
ORF00022	SAG0034	438	sugar ABC transporter, sugar-binding protein
ORF00023	SAG0035	295	sugar ABC transporter, permease protein
ORF00024	SAG0036	276	sugar ABC transporter, permease protein
ORF00025	SAG0037	147	conserved hypothetical protein
ORF00026	SAG0038	220	conserved hypothetical protein
ORF00027	SAG0039	305	N-acetylneuraminate lyase, putative
ORF00028	SAG0040	293	ROK family protein
ORF00029	SAG0041	325	acetyl xylan esterase, putative
ORF00030	SAG0042	267	phosphosugar-binding transcriptional regulator, RpiR family, putative
ORF00031	SAG0043	421	phosphoribosylamine-glycine ligase
ORF00032	SAG0044	162	phosphoribosylaminoimidazole carboxylase, catalytic subunit
ORF00033	SAG0045	363	phosphoribosylaminoimidazole carboxylase, ATPase subunit
ORF00035	SAG0046	463	hypothetical protein
ORF00036	SAG0047	432	adenylosuccinate lyase
ORF00037	SAG0048	303	transcriptional regulator, Cro/CI family
ORF00038	SAG0049	332	Holliday junction DNA helicase RuvB
ORF00039	SAG0050	145	phosphotyrosine protein phosphatase, low molecular weight
ORF00040	SAG0051	126	MORN motif family protein
ORF00041	SAG0052	592	membrane protein, putative
ORF00042	SAG0053	880	aldehyde-alcohol dehydrogenase
ORF00043	SAG0054	338	alcohol dehydrogenase, propanol-preferring
ORF00044	SAG0055	496	threonine synthase
ORF00045	SAG0056	412	MATE efflux family protein
ORF00046	SAG0057	102	ribosomal protein S10
ORF00047	SAG0058	208	ribosomal protein L3
ORF00048	SAG0059	207	ribosomal protein L4
ORF00049	SAG0060	98	ribosomal protein L23
ORF00050	SAG0061	277	ribosomal protein L2
ORF00052	SAG0062	92	ribosomal protein S19
ORF00054	SAG0063	114	ribosomal protein L22
ORF00055	SAG0064	217	ribosomal protein S3
		1	I Protein av





### Table 32: C nv rsi n of ORF Ref Nos. with SAG Ref Nos.

ORF Ref No.	SAGxxxx Ref No.	aa	Annotati n
ORF00056	SAG0065	137	ribosomal protein L16
ORF00058	SAG0066	68	ribosomal protein L29
ORF00059	SAG0067	86	ribosomal protein S17
ORF00060	SAG0068	122	ribosomal protein L14
ORF00061	SAG0069	101	ribosomal protein L24
ORF00063	\$AG0070	180	ribosomal protein L5
ORF00064	SAG0071	61	ribosomal protein S14, putative
ORF00065	SAG0072	132	ribosomal protein S8
ORF00066	SAG0073	178	ribosomal protein L6
ORF00068	SAG0074	118	ribosomal protein L18
ORF00069	SAG0075	164	ribosomal protein S5
ORF00070	SAG0076	59	ribosomal protein L30
ORF00071	SAG0077	146	ribosomal protein L15
ORF00072	SAG0078	434	preprotein translocase, SecY subunit
ORF00073	SAG0079	212	adenylate kinase
ORF00074	SAG0080	72	translation initiation factor IF-1
ORF00075	SAG0081	38	ribosomal protein L36
ORF00077	SAG0082	121	ribosomal protein S13
ORF00078	SAG0083	118	ribosomal protein S11
ORF00080	SAG0084	312	DNA-directed RNA polymerase, alpha subunit
ORF00081	SAG0085	128	ribosomal protein L17
ORF00087	SAG0086	97	hypothetical protein
ORF00088	SAG0087	59	hypothetical protein
ORF00089	SAG0088	56	hypothetical protein
ORF00090	SAG0089	183	conserved hypothetical protein
ORF00091	SAG0090	139	conserved hypothetical protein
ORF00093	SAG0091	144	transcriptional regulator ComX1, putative
ORF00094	SAG0092	230	phosphoglycerate mutase family protein
ORF00095	SAG0093	250	D-alanyl-D-alanine carboxypeptidase family protein
			and the second s
ORF00096	SAG0094	191	N-acetylmuramoyi-L-alanine amidase, family 4 protein
ORF00097	SAG0095	344	heat-inducible transcription repressor HrcA
ORF00098	SAG0096	190	heat shock protein GrpE
ORF00099	SAG0097	609	dnaK protein
ORF00100	SAG0098	379	dnaJ protein
ORF00101	SAG0099	415	transcriptional regulator, GntR family
ORF00102	SAG0100	258	tRNA pseudouridine synthase A
ORF00103	SAG0101	252	phosphomethylpyrimidine kinase, putative
ORF00104	SAG0102	154	conserved hypothetical protein
ORF00105	SAG0103	189	conserved hypothetical protein
ORF00106	SAG0104	280	conserved hypothetical protein
ORF00107	SAG0105	427	trigger factor
ORF00108	SAG0106	191	DNA-directed RNA polymerase, delta subunit, putative
ORF00109	SAG0107	534	CTP synthase
ORF00110	SAG0108	308	conserved hypothetical protein
ORF00111	SAG0109	148	deoxyuridine 5'-triphosphate nucleotidohydrolase
ORF00112	SAG0110	454	DNA repair protein RadA
ORF00113	SAG0111	165	carbonic anhydrase-related protein
ORF00115	SAG0112	439	pyridine nucleotide-disulphide oxidoreductase family
			protein
ORF00116	SAG0113	484	glutamyl-tRNA synthetase
ORF00117	SAG0114	322	ribose ABC transporter, p riplasmic D-ribose-binding
			protein



Table 32: Conversi n of ORF Ref N s. with SAG R f Nos.

ORF Ref N .	SAGxxxx Ref No.	aa	Annotati n
ORF00118	SAG0115	310	ribose ABC transporter, permease protein
ORF00119	SAG0116	492	ribose ABC transporter, ATP-binding protein
ORF00120	SAG0117	132	ribose ABC transporter protein RbsD
ORF00121	SAG0118	303	ribokinase
ORF00122	SAG0119	328	ribose operon repressor RbsR
ORF00123	SAG0120	32	hypothetical protein
ORF00124	SAG0121	362	permease, putative
ORF00125	SAG0122	228	ABC transporter, ATP-binding protein
ORF00126	SAG0123	223	DNA-binding response regulator
ORF00128	SAG0124	356	sensor histidine kinase
ORF00129	SAG0125	396	argininosuccinate synthase
ORF00130	SAG0126	462	argininosuccinate lyase
ORF00131	SAG0127	293	fructose-bisphosphate aldolase
ORF00132	SAG0128	305	L-2-hydroxyisocaproate dehydrogenase
ORF00133	SAG0129	62	ribosomal protein L28
ORF00134	SAG0130	121	conserved hypothetical protein
ORF00135	SAG0131	543	DAK2 domain protein
ORF00136	SAG0132	294	SPFH domain/Band 7 family protein
ORF00137	SAG0133	38	conserved hypothetical protein
ORF00138	SAG0134	96	hypothetical protein
ORF00141	SAG0135	246	amino acid ABC transporter, ATP-binding protein
ORF00142	SAG0136	516	amino acid ABC transporter, amino acid-binding
			protein/permease protein
ORF00143	SAG0137	627	conserved hypothetical protein
ORF00145	SAG0138	279	undecaprenol kinase, putative
ORF00146	SAG0139	251	negative regulator of competence MecA, putative
ORF00148	SAG0140	386	glycosyl transferase, group 4 family protein
ORF00149	SAG0141	256	ABC transporter, ATP-binding protein
ORF00150	SAG0142	420	conserved hypothetical protein
ORF00151	SAG0143	410	selenocysteine lyase
ORF00152	SAG0144	147	NifU family protein
ORF00153	SAG0145	472	conserved hypothetical protein
ORF00154	SAG0148	395	penicillin-binding protein 4, putative
ORF00155	SAG0147	411	D-alanyl-D-alanine carboxypeptidase
ORF00156	SAG0148	551	oligopeptide ABC transporter, substrate binding protein putative
ORF00157	SAG0149	304	oligopeptide ABC transporter, permease protein
ORF00158	SAG0150	343	oligopeptide ABC transporter, permease protein
ORF00160	SAG0151	348	oligopeptide ABC transporter, ATP-binding protein
ORF00161	SAG0152	310	oligopeptide ABC transporter, ATP-binding protein
ORF00166	SAG0153	283	4-diphosphocytidyl-2C-methyl-D-erythritol kinase
ORF00167	SAG0154	147	adc operon repressor AdcR
ORF00168	SAG0155	236	zinc ABC transporter, ATP-binding protein
ORF00169	SAG0156	270	zinc ABC transporter, permease protein
ORF00172	SAG0158	419	tyrosyl-tRNA synthetase
ORF00173	SAG0159	765	penicillin-binding protein 1B, putative
ORF00174	SAG0160	1191	DNA-directed RNA polymerase, beta subunit
ORF00176	SAG0161	1216	
ORF00178	SAG0162	121	conserved hypothetical protein
ORF00179	SAG0163	323	competence protein CgIA
ORF00180	SAG0164	282	competence protein CgIB
		,	
ORF00181	SAG0165	151	conserved hypothetical protein



# Table 32: C nversion of ORF Ref Nos. with SAG K f Nos.

ORF Ref N .	SAGxxxx Ref N .	aa	Annotati n
ORF00183	SAG0167	324	conserved hypothetical protein
ORF00184	SAG0168	397	acetate kinase
ORF00186	SAG0169	68	transcriptional regulator, Cro/CI family
ORF00187	SAG0170	45	hypothetical protein
ORF00188	SAG0171	151	hypothetical protein
ORF00189	SAG0172	221	protease, putative
ORF00190	SAG0173	256	pyrroline-5-carboxylate reductase
ORF00191	SAG0174	355	glutamyl-aminopeptidase
ORF00192	SAG0175	79	hypothetical protein
ORF00193	SAG0176	94	conserved hypothetical protein
ORF00194	SAG0177	107	thioredoxin family protein
ORF00195	SAG0178	208	tRNA binding domain protein
ORF00196	SAG0179	238	conserved hypothetical protein
ORF00198	SAG0180	131	single-strand binding protein
ORF00199	SAG0181	214	hydrolase, haloacid dehalogenase-like family
ORF00200	SAG0182	581	sensor histidine kinase, putative
ORF00201	SAG0183	246	response regulator
ORF00203	SAG0184	151	conserved hypothetical protein
ORF00204	SAG0185	242	membrane protein, putative
ORF00205	SAG0186	36	hypothetical protein
ORF00206	SAG0187	542	oligopeptide ABC transporter, oligopeptide-binding
O111 00200	0,00107	342	protein
ORF00207	SAG0188	325	oligopeptide ABC transporter, permease protein
ORF00208	SAG0189	273	oligopeptide ABC transporter, permease protein
ORF00209	SAG0190	267	peptide ABC transporter, ATP-binding protein
ORF00210	SAG0191	208	peptide ABC transporter, ATP-binding protein
ORF00211	SAG0192	676	PTS system, IIABC components
ORF00212	SAG0193	541	alpha amylase family protein
ORF00214	SAG0194	639	transcriptional antiterminator, BglG family
ORF00216	SAG0195	377	IS1548, transposase
ORF00217	SAG0196	66	conserved domain protein
ORF00218	SAG0197	94	PTS system, IIB component, putative
ORF00219	SAG0198	451	PTS system, IIC component, putative
ORF00220	SAG0199	285	transketolase, N-terminal subunit
ORF00221	SAG0200	309	transketolase, C-terminal subunit
ORF00223	SAG0201	419	oxidoreductase, putative
ORF00224	SAG0202	89	ribosomal protein S15
ORF00225	SAG0203	709	polyribonucleotide nucleotidyltransferase
ORF00226	SAG0204	250	conserved hypothetical protein
ORF00227	SAG0205	194	serine O-acetyltransferase
ORF00228	SAG0206	60	hypothetical protein
ORF00229	SAG0207	447	cysteinyl-tRNA synthetase
ORF00230	SAG0208	128	conserved hypothetical protein
ORF00231	SAG0209	251	RNA methyltransferase, TrmH family, group 3
ORF00232	SAG0210	172	conserved hypothetical protein
ORF00233	SAG0211	286	DegV family protein
ORF00234	SAG0212	32	hypothetical protein
ORF00235	SAG0213	39	hypothetical protein
ORF00236	SAG0214	148	ribosomal protein L13
ORF00237	SAG0215	130	ribosomal protein S9
	SAG0216	33	hypothetical protein
<del></del>		1 33	
ORF00238		204	site-specific recombinace phage interroce family.
<del></del>	SAG0217	384	site-specific recombinase, phage integrase family
ORF00238		384 158	site-specific recombinase, phage integrase family transcriptional regulator, Cro/CI family



### Table 32: Conv rsion f ORF Ref Nos. with SAG R f Nos.

ORF Ref No.	SAGxxxx Ref N .	aa	Ann tati n
ORF00242	SAG0220	92	conserved hypothetical protein
ORF00243	SAG0221	76	hypothetical protein
ORF00244	SAG0222	108	conserved domain protein
ORF00245	SAG0223	209	conserved hypothetical protein, fusion
ORF00246	SAG0224	332	replication initiation protein, putative
ORF00247	SAG0225	144	hypothetical protein
ORF00248	SAG0226	418	recombination protein
ORF00249	SAG0227	156	hypothetical protein
ORF00250	SAG0228	111	conserved hypothetical protein
ORF00251	SAG0229	95	conserved hypothetical protein
ORF00252	SAG0230	96	conserved hypothetical protein
ORF00253	SAG0231	135	hypothetical protein
ORF00254	SAG0232	186	hypothetical protein
ORF00255	SAG0233	226	hypothetical protein
ORF00256	SAG0234	128	hypothetical protein
ORF00257	SAG0235	93	hypothetical protein
ORF00258	SAG0236	32	hypothetical protein
ORF00259	SAG0237	34	hypothetical protein
ORF00260	SAG0238	41	hypothetical protein
ORF00261	SAG0239	286	transcriptional regulator MutR family
ORF00262	SAG0240	393	transporter, putative
ORF00263	SAG0240	213	amino acid ABC transporter, permease protein
ORF00264	SAG0241	308	amino acid ABC transporter, amino acid-binding
OKI-00204	3AG0242	300	protein
ORF00265	SAG0243	211	amino acid ABC transporter, permease protein
ORF00266	SAG0244	381	amino acid ABC transporter, ATP-binding protein
ORF00272	SAG0245	152	hypothetical protein
ORF00273	SAG0246	268	hypothetical protein
ORF00274	SAG0247	116	hypothetical protein
ORF00275	SAG0248	90	hypothetical protein
ORF00276	SAG0249	116	hypothetical protein
ORF00278	SAG0250	193	hypothetical protein
ORF00279	SAG0251	72	transcriptional regulator, Cro/CI family
ORF00280	SAG0252	186	acetyltransferase, GNAT family
ORF00281	SAG0253	192	acetyltransferase, GNAT family
ORF00282	SAG0254	226	acetyltransferase, GNAT family
ORF00283	SAG0255	315	conserved hypothetical protein
ORF00284	SAG0256	163	RNA polymerase sigma factor, ECF subfamily
ORF00285	SAG0257	53	hypothetical protein
ORF00287	SAG0258	202	transcriptional regulator, TetR family
ORF00288	SAG0259	365	ABC transporter efflux protein, DrrB family, putative
			7,
ORF00289	SAG0260	238	ABC transporter, ATP-binding protein
ORF00290	SAG0261	129	IS1381, transposase OrfB
ORF00291	SAG0262	127	IS1381, transposase OrfA
ORF00292	SAG0263	171	hypothetical protein
ORF00293	SAG0264	103	conserved hypothetical protein
ORF00294	SAG0265	235	conserved hypothetical protein
ORF00295	SAG0266	382	N-acetylglucosamine-6-phosphate deacetylase
ORF00296	SAG0267	180	conserved hypothetical protein
ORF00297	SAG0268	304	glycyl-tRNA synthetase, alpha subunit
ORF00298	SAG0269	213	acyl carrier protein phosphodiesterase, putative
ORF00299	SAG0270	679	glycyl-tRNA synthetase, beta subunit
ORF00300	SAG0271	85	conserved hypothetical protein
ORF00301	SAG0272	87	membrane protein, putativ





### Tabl 32: Conv rsion of ORF Ref Nos. with SAG R f Nos.

ORF Ref N .	SAGxxxx Ref No.	aa	Annotation
ORF00302	SAG0273	502	glycerol kinase
ORF00303	SAG0274	609	alpha-glycerophosphate oxidase
ORF00304	SAG0275	232	glycerol uptake facilitator protein
ORF00305	SAG0276	445	NADH oxidase, putative
ORF00306	SAG0277	476	conserved hypothetical protein
ORF00307	SAG0278	661	transketolase
ORF00308	SAG0279	101	conserved hypothetical protein
ORF00309	SAG0280	244	ABC transporter, ATP-binding protein
ORF00310	SAG0281	534	membrane protein, putative
ORF00313	SAG0282	461	PTS system, IIBC components
ORF00314	SAG0283	267	glutamate 5-kinase
ORF00315	SAG0284	417	gamma-glutamyl phosphate reductase
ORF00316	SAG0285	298	conserved hypothetical protein TIGR00006
ORF00317	SAG0286	108	cell division protein FtsL, putative
ORF00318	SAG0287	752	penicillin-binding protein 2X
ORF00319	SAG0288	336	phospho-N-acetylmuramoyl-pentapeptide-transferase
			prospire is addynialaneys portapopado danoicidad
ORF00320	SAG0289	447	ATP-dependent RNA helicase, DEAD/DEAH box
			family
ORF00321	SAG0290	270	ABC transporter, substrate-binding protein
. ORF00322	SAG0291	267	amino acid ABC transporter, permease protein
ORF00323	SAG0292	247	amino acid ABC transporter, ATP-binding protein
ORF00324	SAG0293	74	conserved hypothetical protein
ORF00325	SAG0294	304	thioredoxin reductase
ORF00326	SAG0295	486	conserved hypothetical protein
ORF00327	SAG0296	273	NAD synthetase
ORF00328	SAG0297	444	aminopeptidase C
ORF00329	SAG0298	750	penicillin-binding protein 1A
ORF00330	SAG0299	199	recombination protein U
ORF00331	SAG0300	172	conserved hypothetical protein
ORF00332	SAG0301	40	hypothetical protein
ORF00333	SAG0302	110	conserved hypothetical protein
ORF00335	SAG0303	384	conserved hypothetical protein
ORF00336	SAG0304	487	conserved hypothetical protein
ORF00337	SAG0305	160	autoinducer-2 production protein LuxS
ORF00338	SAG0306	535	KH domain protein
ORF00340	SAG0307	33	hypothetical protein
ORF00341	SAG0308		ABC transporter, ATP-binding protein, FRAMESHIFT
ORF00343	SAG0309	246	ABC transporter, permease protein, putative
ORF00344	SAG0310	361	conserved hypothetical protein
ORF00345	SAG0311		DNA-binding response regulator POINT MUTATION
			,
ORF00347	SAG0312	234	conserved hypothetical protein
ORF00348	SAG0313	209	guanylate kinase
ORF00349	SAG0314	104	DNA-directed RNA polymerase, omega subunit,
0000000		ļ	putative
ORF00350	SAG0315	796	primosomal protein N'
ORF00351	SAG0316	311	methionyl-tRNA formyltransferase
ORF00352	SAG0317	440	Sun protein
ORF00353	SAG0318	245	serine/threonine phosphatase, putative
ORF00354	SAG0319	651	serine/threonine protein kinase
ORF00355	SAG0320	231	conserved hypothetical prot in
ORF00356	SAG0321	339	sensor histidin kinase, putative
ORF00358	SAG0322	213	DNA-binding response regulator

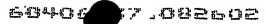


# Table 32: Conv rsi n of ORF Ref Nos. with SAGR f Nos.

ORFR f No.	SAGxxxx Ref No.	aa	Ann tati n
ORF00359	SAG0323	466	hydrolase, haloacid dehalogenase family/peptidyl-
			prolyl cis-trans isomerase, cyclophilin type
ORF00360	SAG0324	124	general stress protein, putative
ORF00361	SAG0325	258	pyruvate formate-lyase-activating enzyme
ORF00362	SAG0326	251	transcriptional regulator, DeoR family
ORF00363	SAG0327	327	transcriptional regulator, putative
ORF00364	SAG0328	107	PTS system, cellobiose-specific IIA component
ORF00366	SAG0329	106	PTS system, cellobiose-specific IIB component
ORF00367	SAG0330	433	PTS system, cellobiose-specific IIC component
ORF00368	SAG0331	818	formate acetyltransferase
ORF00369	SAG0332	222	transaldolase family protein
ORF00371	SAG0333	362	glycerol dehydrogenase
ORF00372	SAG0334	308	cysteine synthase A
ORF00373	SAG0335	214	
ORF00374			conserved hypothetical protein TIGR00257
	SAG0336	429	helicase, putative
ORF00375	SAG0337	221	competence protein F, putative
ORF00376	SAG0338	184	ribosomal subunit interface protein
ORF00382	SAG0339	450	aspartate kinase family protein
ORF00383	SAG0340	216	hydrolase, haloacid dehalogenase-like family
ORF00384	SAG0341	49	hypothetical protein
ORF00385	SAG0342	263	enoyl-CoA hydratase/isomerase family protein
ORF00386	SAG0343	144	transcriptional regulator, MarR family
ORF00387	SAG0344	323	3-oxoacyl-(acyl-carrier-protein) synthase III
ORF00388	SAG0345	74	acyl carrier protein
ORF00390	SAG0346	319	enoyl-(acyl-carrier-protein) reductase II
ORF00391	SAG0347	308	malonyl CoA-acyl carrier protein transacylase
ORF00392	SAG034B	244	3-oxoacyl-[acyl-carrier protein] reductase
ORF00393	SAG0349	410	3-oxoacyl-(acyl-carrier-protein) synthase II
ORF00394	SAG0350	166	acetyl-CoA carboxylase, biotin carboxyl carrier protein
ORF00395	SAG0351	140	(3R)-hydroxymyristoyl-(acyl-carrier-protein) dehydratase
ORF00396	SAG0352	456	acetyl-CoA carboxylase, biotin carboxylase
ORF00397	SAG0353	291	acetyl-CoA carboxylase, carboxyl transferase, beta subunit
ORF00398	SAG0354	257	acetyl-CoA carboxylase, carboxyl transferase, alpha subunit
ORF00399	SAG0355	210	conserved hypothetical protein
ORF00400	SAG0356	425	seryl-tRNA synthetase
ORF00402	SAG0357	330	hypothetical protein
ORF00403	SAG0358	120	conserved hypothetical protein
ORF00404	SAG0359	303	PTS system, mannose-specific IID component
ORF00405	SAG0360	270	PTS system, mannose-specific IIC component
ORF00406	SAG0361	336	PTS system, mannose-specific IIAB components
ORF00407	SAG0362	270	hydrolase, haloacid dehalogenase-like family
ORF00408	SAG0363	194	hypothetical protein .
ORF00409	SAG0364	203	membrane protein, putative
ORF00410	SAG0365	473	xanthine/uracil permease family protein
ORF00411	SAG0366	169	conserved hypothetical protein TIGR00150, putative
ORF00412	SAG0367	186	acetyltransferase, GNAT family
ORF00413	SAG0368	435	transcriptional regulator, putative
ORF00414	SAG0369	98	conserved hypothetical protein
ORF00415	SAG0370	139	HIT family protein
ORF00416	SAG0371	167	hypothetical protein
ORF00417	SAG0372	85	hypoth tical protein



ORF Ref No.	SAGxxxx Ref No.	aa	Ann tati n
ORF00419	SAG0373	241	ABC transporter, ATP-binding protein
ORF00421	SAG0374	344	ABC transporter, permease protein
ORF00422	SAG0375	266	conserved hypothetical protein
ORF00423	SAG0376	211	conserved hypothetical protein TIGR00091
ORF00424	SAG0377	127	conserved hypothetical protein, POINT MUTATION
		, ,_,	Proton, Political Proton, Political Proton
ORF00425	SAG0378	379	N utilization substance protein A
ORF00426	SAG0379	98	conserved hypothetical protein
ORF00427	SAG0380	100	ribosomal protein L7A family
ORF00428	SAG0381	927	translation initiation factor IF-2
ORF00429	SAG0382	122	ribosome-binding factor A
ORF00430	SAG0383	334	conserved hypothetical protein
ORF00431	SAG0384	138	transcriptional repressor CopY
ORF00432	SAG0385	744	copper-transporter ATPase CopA
ORF00433	SAG0386	68	copper-transporter protein CopZ
ORF00434	SAG0387	204	conserved hypothetical protein
ORF00435	SAG0388	270	hydrolase, haloacid dehalogenase-like family
ORF00436	SAG0389	880	DNA polymerase I
ORF00437	SAG0390	146	CoA binding domain protein
ORF00438	SAG0391	159	transcriptional regulator, Fur family
ORF00439	SAG0392	521	cell wall surface anchor family protein
ORF00440	SAG0393	228	DNA-binding response regulator
ORF00441	SAG0394	345	sensor histidine kinase
ORF00442	SAG0395	246	conserved hypothetical protein
ORF00443	SAG0396	380	queuine tRNA-ribosyltransferase
ORF00444	SAG0397	102	conserved hypothetical protein
ORF00445	SAG0398	179	bioY family protein
ORF00446	SAG0399	258	AtsA/ElaC family protein
ORF00447	SAG0400	168	cytidine/deoxycytidylate deaminase family protein
ORF00448	SAG0401	44	hypothetical protein
ORF00449	SAG0402	449	glucose-6-phosphate isomerase
ORF00450	SAG0403	175	5-formyltetrahydrofolate cyclo-ligase family protein
ORF00451	SAG0404	225	showhold for the protein
ORF00452	SAG0404 SAG0405	347	rhomboid family protein
ORF00453	SAG0406	299	UTP-glucose-1-phosphate uridylyltransferase
ORF00454	SAG0400 SAG0407	338	glycerol-3-phosphate dehydrogenase (NAD(P)+)
ORF00455	SAG0408	109	ribonuclease P protein component
ORF00456	SAG0409	271	SpollIJ family protein
ORF00458	SAG0410	273	R3H domain protein
ORF00463	SAG0411	177	conserved hypothetical protein
ORF00464	SAG0412	258	RecX protein
ORF00465	SAG0412	451	RNA methyltransferase, TrmA family
ORF00466	SAG0414	153	conserved hypothetical protein
ORF00467	SAG0414 SAG0415	142	acetyltransferase, GNAT family
ORF00468	SAG0416	1233	protease, putative
ORF00469	SAG0417	302	glycosyl transferase, group 2 family protein
ORF00470	SAG0418	336	ribonucleoside-diphosphate reductase 2, beta subunit
2.0.0000	37.00410	~~	managed and complication reductase 2, Deta Subunit
ORF00471	SAG0419	137	nrdl protein
ORF00472	SAG0420	721	ribonucleoside-diphosphate reductase 2, alpha subun
ORF00473	SAG0421	1055	conserved hypothetical protein
ORF00474	SAG0422	129	conserved hypothetical protein
ORF00475	SAG0423	132	conserved domain protein
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### Tabl 32: Conv rsi n of ORF Ref Nos. with SAGR f Nos.

ORF Ref N .	SAGxxxx Ref N .	aa	Ann tation
ORF00476	SAG0424	94	hypothetical protein .
ORF00478	SAG0425	105	carboxymuconolactone decarboxylase family protein
ORF00479	SAG0426	131	conserved hypothetical protein
ORF00480	SAG0427	129	transcriptional regulator, MerR family
ORF00482	SAG0428	345	alcohol dehydrogenase, zinc-containing
ORF00483	SAG0429	284	oxidoreductase, aldo/keto reductase family
ORF00484	SAG0430	287	cation efflux system protein
ORF00485	SAG0431	174	transcriptional regulator, TetR family
ORF00486	SAG0432	397	transcriptional regulator, AraC family
ORF00487	SAG0433	1389	surface protein Rib
ORF00488	SAG0434	61	transposase, IS256 family, truncation
ORF00489	SAG0435	97	DNA-damage-inducible protein J, putative
ORF00490	SAG0436	62	hypothetical protein
ORF00491	SAG0437	123	hypothetical protein
ORF00493	SAG0438	145	bacteriophage L54a, integrase, truncation
ORF00495	SAG0439		conserved hypothetical protein, FRAMESHIFT
ORF00496	SAG0440	84	conserved hypothetical protein
ORF00497	SAG0441	103	conserved domain protein
ORF00499	SAG0442	189	acetyltransferase, GNAT family
ORF00500	SAG0443	194	acetyltransferase, GNAT family
ORF00501	SAG0444	188	conserved hypothetical protein
ORF00502	SAG0445	883	valyI-tRNA synthetase
ORF00503	SAG0446 ·	319	oxidoreductase, Gfo/Idh/MocA family
ORF00504	SAG0447	287	magnesium transporter, CorA family
ORF00506	SAG0448	391	transposase, IS256 family
ORF00507	SAG0449	354	conserved hypothetical protein
ORF00508	SAG0450	330	aspartate-ammonia ligase
ORF00510	SAG0451	149	bacteriocin transport accessory protein, putative
ORF00511	SAG0452	179	type II DNA modification methyltransferase, putative
ORF00512	SAG0453	96	hypothetical protein
ORF00513	SAG0454	161	phosphopantetheine adenylyltransferase
ORF00515	SAG0455	357	conserved hypothetical protein
ORF00518	SAG0456		conserved hypothetical protein, degenerate
ORF00519	SAG0457	192	conserved hypothetical protein
ORF00520	SAG0458	368	conserved hypothetical protein TIGR00048
ORF00521	SAG0459	171	VanZF domain protein
ORF00522	SAG0460	581	ABC transporter, ATP-binding/permease protein
ORF00523	SAG0461	579	ABC transporter, ATP-binding/permease protein
ORF00524	SAG0462	188	anthranilate synthase component II
ORF00525	SAG0463	179	bioY family protein
ORF00526	SAG0464	330	biotin synthetase
ORF00527	SAG0465	164	hypothetical protein
ORF00528	SAG0466	371	thiolase
ORF00531	SAG0467	409	AMP-binding enzyme domain protein
ORF00532	SAG0468	210	endonuclease III
ORF00533	SAG0469	131	type IV prepilin peptidase-related protein
ORF00534	SAG0470	69	conserved hypothetical protein
ORF00535	SAG0471	322	glucokinase
ORF00536	SAG0472	126	rhodanese domain protein
ORF00537	SAG0473	613	elongation factor Tu family protein
ORF00538	SAG0474	81	conserved hypothetical protein
ORF00540	SAG0475	451	UDP-N-acetylmuramoylalanine—D-glutamate ligase
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Table 32: Conversi n of ORF Ref Nos. with SAGR f Nos.

ORFR fNo.	SAGxxxx Ref No.	aa	Annotation
ORF00541	SAG0476	358	UDP-N-acetylglucosamineN-acetylmuramyl-
			(pentapeptide) pyrophosphoryl-undecaprenol N-
	0.00.		acetylglucosamine transferase
ORF00542	SAG0477	378	cell division protein DivIB, putative
ORF00544	SAG0478	429	cell division protein FtsA
ORF00545	SAG0479	426	cell division protein FtsZ
ORF00546	SAG0480	224	ylmE protein, putative
ORF00547	SAG0481	201	ylmF protein
ORF00548	SAG0482	84	YGGT family protein
ORF00549	SAG0483	262	ylmH protein
ORF00550	SAG0484	256	cell division protein DivIVA, putative
ORF00552	SAG0485	930	isoleucyl-tRNA synthetase
ORF00553	SAG0486	100	conserved hypothetical protein
ORF00554	SAG0487	151	MutT/nudix family protein
ORF00555	SAG0488	753	ATP-dependent Clp protease, ATP-binding subunit
ORF00556	SAG0489	34	hypothetical protein
ORF00557	SAG0490	76	conserved hypothetical protein
ORF00558	SAG0491	230	amino acid ABC transporter, permease protein
ORF00559	SAG0492	244	amino acid ABC transporter, ATP-binding protein
ORF00560	SAG0493	564	phosphoglucomutase/phosphomannomutase family protein
ORF00562	SAG0494	284	methylenetetrahydrofolate
		ļ	dehydrogenase/methenyltetrahydrofolate
		<u> </u>	cyclohydrolase
ORF00563	SAG0495	278	conserved hypothetical protein
ORF00564	SAG0496	446	exodeoxyribonuclease VII, large subunit
ORF00565	SAG0497	71	exodeoxyribonuclease VII, small subunit
ORF00566	SAG0498	290	geranyltranstransferase, putative
ORF00567	SAG0499	275	hemolysin A
ORF00568	SAG0500	157	arginine repressor ArgR, putative
ORF00570	SAG0501	552	DNA repair protein RecN
ORF00571	SAG0502	278	DegV family protein
ORF00572	SAG0503	279	Lipase/Acylhydrolase, putative
ORF00573	SAG0504	200	conserved hypothetical protein
ORF00574	SAG0505	91	DNA-binding protein HU
ORF00575	SAG0506	65	hypothetical protein
ORF00576	SAG0507	310	dihydroorotate dehydrogenase A
ORF00577	SAG0508	411	beta-lactam resistance factor
ORF00578	SAG0509	403	beta-lactam resistance factor
ORF00579	SAG0510	406	murM protein, putative
ORF00580	SAG0511	270	hydrolase, haloacid dehalogenase-like family
ORF00581	SAG0512	438	HD domain protein
ORF00582	SAG0513	128	conserved hypothetical protein
ORF00583	SAG0514	894	cation-transporting ATPase, E1-E2 family
ORF00584	SAG0515	286	conserved hypothetical protein
ORF00585	SAG0516	643	fructose-1,6-bisphosphatase, putative
ORF00586	SAG0517	374	iron-sulfur cluster-binding protein, putative
ORF00587	SAG0518		peptide chain release factor 2, FRAMESHIFT
ORF00588	SAG0519	230	cell division ABC transport r, ATP-binding protein Fi
ORF00589	SAG0520	309	cell division ABC transporter, permease protein FtsX
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ORF00590	SAG0521	236	carboxymethylenebut nolidase-related protein





### Table 32: Conv rsion f ORF Ref Nos. with SAG K f Nos.

ORFR f No.	SAGxxxx Ref No.	aa	Ann tation
ORF00592	SAG0523	254	oxidoreductase, short chain dehydrogenase/reductase
			family
ORF00593	SAG0524	835	DNA polymerase III, epsilon subunit/ATP-dependent
ODEOGEOG	0400505	007	helicase DinG
ORF00595	SAG0525	397	aspartate aminotransferase
ORF00596	SAG0526	448	asparaginyl-tRNA synthetase
ORF00597	SAG0527	185	conserved hypothetical protein
ORF00598	SAG0528	327	inosine-uridine preferring nucleoside hydrolase
ORF00599	SAG0529	38	hypothetical protein
ORF00600	SAG0530	137	OsmC/Ohr family protein
ORF00601	SAG0531	296	conserved hypothetical protein
ORF00602	SAG0532	324	conserved hypothetical protein
ORF00603	SAG0533	303	Uncharacterized BCR, COG1481
ORF00604	SAG0534	465	dipeptidase
ORF00605	SAG0535	506	zinc ABC transporter, zinc-binding adhesion liprotein
ORF00606	SAG0536	86	ribosomal protein L31
ORF00607	SAG0537	311	DHH family protein
ORF00608	SAG0538	340	adenosine deaminase, putative
ORF00609	SAG0539	147	flavodoxin
ORF00610	SAG0540	91	chorismate mutase, putative
ORF00611	SAG0541	398	voltage-gated chloride channel family protein
ORF00612	SAG0542	127	IS1381, transposase OrfA
ORF00613	SAG0543	129	IS1381, transposase OrfB
ORF00614	SAG0544	115	ribosomal protein L19
ORF00615	SAG0545	359	site-specific recombinase, phage integrase family
			one openio recombinates, priage magrace ranning
ORF00617	SAG0546	67	conserved domain protein
ORF00618	SAG0547	185	hypothetical protein
ORF00619	SAG0548	265	repressor protein, putative
ORF00620	SAG0549	47	hypothetical protein
ORF00621	SAG0550	74	conserved hypothetical protein
ORF00622	SAG0551	52	conserved hypothetical protein
ORF00623	SAG0552	62	hypothetical protein
ORF00624	SAG0553	268	hypothetical protein
ORF00626	SAG0554	63	transcriptional regulator, Cro/Cl family
ORF00627	SAG0555	249	antirepressor, putative
ORF00628	SAG0556	47	hypothetical protein
ORF00630	SAG0557	76	hypothetical protein
ORF00632	SAG0558	74	hypothetical protein
ORF00633	SAG0559	286	conserved hypothetical protein
ORF00634	SAG0560	77	conserved hypothetical protein
ORF00635	SAG0561	46	hypothetical protein
ORF00636	SAG0562	84	hypothetical protein
ORF00637	SAG0563	53	
ORF00638	<del></del>		hypothetical protein
ORF00639	SAG0564	160	conserved hypothetical protein
	SAG0565	224	conserved domain protein
ORF00640	SAG0566	138	single-strand binding protein
ORF00641	SAG0567	439	reverse transcriptase/maturase family protein
ORF00642	SAG0568	67	conserved hypothetical protein
ORF00643	SAG0569	158	conserved hypothetical protein
ORF00644	SAG0570	115	hypothetical protein
ORF00645	SAG0571	43	hypothetical protein
ORF00646	SAG0572	138	conserv d hypothetical protein
ORF00647	SAG0573	54	hypothetical protein

# Tabl 32: Conversion of ORF R f N s. with SAG Ref Nos.

ORF Ref No.	SAGxxxx Ref No.	aa	Annotati n
ORF00648	SAG0574	89	conserved hypothetical protein
ORF00649	SAG0575	110	hypothetical protein
ORF00650	SAG0576	43	hypothetical protein
ORF00652	SAG0577	177	conserved hypothetical protein
ORF00653	SAG0578	88	conserved hypothetical protein
ORF00654	SAG0581	118	conserved hypothetical protein
ORF00655	SAG0582	422	conserved hypothetical protein
ORF00656	SAG0583	406	conserved hypothetical protein
ORF00657	SAG0584	62	conserved hypothetical protein, truncation
ORF00658	SAG0585	471	conserved hypothetical protein
ORF00659	SAG0586	154	conserved hypothetical protein
ORF00660	SAG0587	300	structural protein, putative
ORF00661	SAG0588	71	conserved hypothetical protein
ORF00662	SAG0589	143	conserved hypothetical protein
ORF00663	SAG0590	112	conserved hypothetical protein
ORF00684	SAG0591	78	conserved hypothetical protein
ORF00665	SAG0592	111	
ORF00666	SAG0592 SAG0593	185	conserved hypothetical protein
ORF00667			structural protein
ORF00668	SAG0594	81	conserved hypothetical protein
	SAG0595	123	conserved hypothetical protein
ORF00669	SAG0596	670	PblA, internal deletion
ORF00670	SAG0597	506	minor structural protein, putative
ORF00671	SAG0598	1374	minor structural protein, putative
ORF00672	SAG0599	668	minor structural protein, putative
ORF00673	SAG0600	109	hypothetical protein
ORF00674	SAG0601	70.	hypothetical protein
ORF00675	SAG0602	100	conserved hypothetical protein
ORF00676	SAG0603	111	holin, putative
ORF00677	SAG0604	239	lysin, putative
ORF00678	SAG0605	323	conserved hypothetical protein
ORF00679	SAG0606	66	conserved hypothetical protein
ORF00681	SAG0607	56	conserved hypothetical protein
ORF00682	SAG0608	59	hypothetical protein
ORF00683	SAG0609	193	site-specific recombinase, phage integrase family
ORF00685	SAG0610	134	conserved hypothetical protein
ORF00687	SAG0611		transposase, degenerate FRAMESHIFT
ORF00689	SAG0612	53	conserved hypothetical protein, FRAMESHIFT
ORF00690	SAG0613	425	transmembrane protein Vexp1
ORF00691	SAG0614	218	ABC transporter, ATP-binding protein Vexp2
ORF00692	SAG0615	458	transmembrane protein Vexp3
ORF00693	SAG0616	217	DNA-binding response regulator VncR
ORF00694	SAG0617	439	sensor histidine kinase VncS
ORF00695	SAG0618	195	transposase OrfB, IS3 family, truncation
ORF00697	SAG0619	66	conserved hypothetical protein
ORF00698	SAG0620	62	hypothetical protein
ORF00699	SAG0621	401	rod shape-determining protein RodA, putative
ORF00700	SAG0622	186	hydrolase, haloacid dehalogenase-like family
ORF00701	SAG0623	650	DNA gyrase, B subunit
ORF00702	SAG0624	574	septation ring formation regulator EzrA, putative
ORF00703	SAG0625	213	phosphoserine phosphatase SerB
ORF00704	SAG0626	161	MutT/nudix family protein
ORF00705	SAG0627	151	conserved hypothetical protein
ORF00706	SAG0628	435	
ORF00707	SAG0629		enolase
	0/10/029	354	conserved domain protein





# Table 32: C nversion f ORF R f Nos. with SAG R f Nos.

ORF Ref No.	SAGxxxx Ref No.	aa	Annotation
ORF00708	SAG0630	427	3-phosphoshikimate 1-carboxyvinyltransferase
ORF00709	SAG0631	170	shikimate kinase
ORF00710	SAG0632	457	psr protein
ORF00711	SAG0633	451	RNA methyltransferase, TrmA family
ORF00712	SAG0634	70	hypothetical protein
ORF00713	SAG0635	245	acid phosphatase precursor, class B
ORF00714	SAG0636	172	conserved hypothetical protein
ORF00717	SAG0637		transcriptional regulator, TetR family, putative,
]	G. 150007		FRAMESHIFT
ORF00718	SAG0638	109	cell wall surface anchor family protein
ORF00720	SAG0639	273	transposase OrfB, IS3 family
ORF00721	SAG0640	91	transposase OrfA, IS3 family
ORF00722	SAG0641		Tn5252, Orf 10 protein, degenerate POINT MUTATION
		1	mozoz, on to protein, degenerate t onth motation
ORF00723	SAG0642	59	hypothetical protein
ORF00725	SAG0843		chaperonin, 33 kDa DEGENERATE
ORF00726	SAG0644	402	transcriptional regulator, AraC family
ORF00727	SAG0645	554	cell wall surface anchor family protein, putative
ORF00728	SAG0646	307	cell wall surface anchor family protein
ORF00729	SAG0647	305	sortase family protein
ORF00731	SAG0648	260	sortase family protein
ORF00732	SAG0649	890	cell wall surface anchor family protein putative
ORF00734	SAG0650	189	sortase family protein, FRAMESHIFT
ORF00735	SAG0651	201	hypothetical protein
ORF00737	SAG0653	76	conserved hypothetical protein, DEGENERATE
ORF00738	SAG0654	34	hypothetical protein
ORF00740	SAG0656	36	hypothetical protein
ORF00741	SAG0657	89	hypothetical protein
ORF00742	SAG0658	383	lipoprotein, putative
ORF00743	SAG0659	330	ABC transporter, ATP-binding protein
ORF00744	SAG0660	272	membrane protein
ORF00745	SAG0661	261	conserved hypothetical protein
ORF00747	SAG0663	282	cyID protein
ORF00748	SAG0664	240	cylG protein
ORF00749	SAG0665	101	acyl carrier protein AcpC
ORF00750	SAG0666	158	cylZ protein FRAMESHIFT
ORF00751	SAG0667	309	cylA protein
ORF00752	SAG0668	292	cylB protein
ORF00753	SAG0669	667	cylE protein
ORF00754	SAG0670	317	cylF protein
ORF00755	SAG0671	731	cyll protein
ORF00756	SAG0672	403	cylJ protein
ORF00757	SAG0673	191	cylK protein
ORF00758	SAG0674	113	hypothetical protein
ORF00759	SAG0675	171	surface protein antigen-related protein
ORF00760	SAG0676	885	serine protease, subtilase family, putative
ORF00761	SAG0677	1062	hypothetical protein
ORF00762	SAG0678	T	endopeptidase O DEGENERATE
ORF00766	SAG0679	286	hydrolase, alpha/beta fold family, putative
ORF00767	SAG0680	339	hypothetical protein
ORF00768	SAG0681	353	conserved domain protein
ORF00769	SAG0682	409	permease, putative
ORF00770	SAG0683		transmembrane protein Vexp3, putative FRAMESHIFT
ORF00774	SAG0684	223	ABC transporter, ATP-binding protein



### Table 32: C nv rsi n of ORF Ref Nos. with SAG R f Nos.

ORF Ref N .	SAGxxxx Ref No.	aa	Annotation
ORF00775	SAG0685	472	conserved hypothetical protein
ORF00776	SAG0686	261	DNA-entry nuclease, putative
ORF00777	SAG0687	212	DedA family protein, putative
ORF00778	SAG0688	218	ABC transporter, ATP-binding protein
ORF00779	SAG0689	257	membrane protein, putative
ORF00780	SAG0690	272	conserved hypothetical protein
ORF00781	SAG0691	294	transcriptional regulator, LysR family
ORF00783	SAG0692	193	regulatory protein, putative
ORF00785	SAG0693	377	IS1548, transposase
ORF00786	SAG0694	173	regulatory protein, putative, truncation
ORF00787	SAG0695	330	D-lactate dehydrogenase
ORF00788	SAG0696	516	sodium:galactoside symporter family protein, putative
J 30733	0,10000	0,0	Souldinguisologide Sympotics lathing protein, putable
ORF00789	SAG0697	341	2-keto-3-deoxygluconate kinase
ORF00790	SAG0698	599	beta-glucuronidase
ORF00791	SAG0699	223	transcriptional regulator, GntR family
ORF00792	SAG0700	205	2-dehydro-3-deoxyphosphogluconate aldolase/4-
0711 00702	07100100	200	hydroxy-2-oxoglutarate aldolase
ORF00793	SAG0701	466	Glucuronate isomerase
ORF00794	SAG0702	348	mannonate dehydratase
ORF00795	SAG0703	279	D-mannonate oxidoreductase
ORF00796	SAG0704	270	hydrolase, haloacid dehalogenase-like family
ORF00797	SAG0705	596	glycosyl hydrolase, family 3
ORF00798	SAG0705	361	proline dipeptidase
ORF00799	SAG0708	<del></del>	<u> </u>
ORF00800		334	transcriptional regulator, RegM family
	SAG0708	488	alpha amylase family protein
ORF00801	SAG0709	332	glycosyl transferase, group 1 family protein
ORF00802	SAG0710	444	glycosyl transferase, group 1 family protein
ORF00803	SAG0711	647	threonyl-tRNA synthetase
ORF00804	SAG0712	234	DNA-binding response regulator
ORF00805	SAG0713	339	conserved hypothetical protein
ORF00806	SAG0714	188	conserved hypothetical protein
ORF00807	SAG0715	216	amino acid ABC transporter, permease protein
ORF00808	SAG0716	231	amino acid ABC transporter, permease protein
ORF00809	SAG0717	266	amino acid ABC transporter, amino acid-binding
0000000	0100740	- 054	protein
ORF00810	SAG0718	251	amino acid ABC transporter, ATP-binding protein
ORF00811	SAG0719	236	DNA-binding response regulator
ORF00812	SAG0720	449	sensory box histidine kinase
ORF00813	SAG0721	269	metallo-beta-lactamase family protein
ORF00814	SAG0722	122	conserved hypothetical protein
ORF00815	SAG0723	236	ribonuclease III
ORF00816	SAG0724	1179	SMC family protein
ORF00817	SAG0725	265	hydrolase, haloacid dehalogenase-like family
ORF00818	SAG0726	274	hydrolase, haloacid dehalogenase-like family
ORF00819	SAG0727	536	signal recognition particle-docking protein FtsY
ORF00820	SAG0728	270	ABC transporter, substrate-binding protein
ORF00821	SAG0729	300	ABC transporter, permease protein, putative
ORF00822	SAG0730	42	ABC transporter, ATP-binding protein
ORF00823	SAG0731	347	bacterial luciferase family protein
ORF00824	SAG0732	720	transcriptional accessory protein Tex, putative
ORF00825	SAG0733	142	conserved hypothetical protein
ORF00826	SAG0734	87	phage shock protein C, putativ
ORF00827	SAG0735	44	hypothetical protein
ORF00828	SAG0736	311	HPr(Ser) kinase/phosphatase
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ORF Ref No.	SAGxxxx Ref No.	aa	Ann tation
ORF00830	SAG0737	257	prolipoprotein diacylglyceryl transferase
ORF00832	SAG0738	132	conserved hypothetical protein
ORF00833	SAG0739	143	conserved hypothetical protein
ORF00834	SAG0740	91	conserved hypothetical protein
ORF00835	SAG0741	303	peptidase, U32 family, putative
ORF00836	SAG0742	428	peptidase, U32 family
ORF00837	SAG0743	70	conserved hypothetical protein
ORF00838	SAG0744	265	membrane protein, putative
ORF00839	SAG0745	446	Mn2+/Fe2+ transporter, NRAMP family
ORF00840	SAG0746	369	riboflavin biosynthesis protein RibD
ORF00841	SAG0747	208	riboflavin synthase, alpha subunit
ORF00842	SAG0748	397	riboflavin biosynthesis protein RibA
ORF00843	SAG0749	156	riboflavin synthase, beta subunit
ORF00844	SAG0750	496	lysyl-tRNA synthetase
ORF00845	SAG0751	300	hydrolase, haloacid dehalogenase-like family
ORF00846	SAG0752	213	phosphoglycerate mutase family protein
ORF00847	SAG0753	157	ebsC family protein, putative
ORF00848	SAG0754	205	conserved domain protein
ORF00850	SAG0755	282	peptidase, U32 family
ORF00852	SAG0756	174	conserved hypothetical protein
ORF00853	SAG0757	129	lipoprotein, putative
ORF00855	SAG0758	599	oligoendopeptidase F, putative
ORF00856	SAG0759	931	phosphoenolpyruvate carboxylase
ORF00857	SAG0760	377	IS1548, transposase
ORF00859	SAG0761	422	cell division protein, FtsW/RodA/SpoVE family
ORF00861	SAG0762	398	translation elongation factor Tu
ORF00863	SAG0763	252	triosephosphate isomerase
ORF00865	SAG0764	230	phosphoglycerate mutase
ORF00866	SAG0765	681	penicillin-binding protein 2b
ORF00867	SAG0766	198	recombination protein RecR
ORF00868	SAG0767	348	D-alanineD-alanine ligase
ORF00869	SAG0768	455	UDP-N-acetylmuramoylalanyl-D-glutamyl-2,6-
			diaminopimelateD-alanyl-D-alanyl ligase
ORF00870	SAG0769	406	oxalate:formate antiporter
ORF00871	SAG0770	228	conserved hypothetical protein
ORF00872	SAG0771	512	cell wall surface anchor family protein
ORF00873	SAG0772	514	peptide chain release factor 3
ORF00874	SAG0773	126	conserved hypothetical protein
ORF00876	SAG0774	244	ABC transporter, ATP-binding protein
ORF00878	SAG0775	220	ABC transporter, permease protein
ORF00879	SAG0776	276	lipoprotein, putative
ORF00880	SAG0777	528	ATP-dependent RNA helicase, DEAD/DEAH box
	····		family
ORF00882	SAG0778	88	conserved hypothetical protein
ORF00883	SAG0779	254	conserved hypothetical protein
ORF00884	SAG0780	246	acyltransferase family protein
ORF00885	SAG0781	217	competence protein CelA
ORF00887	SAG0782	745	DNA internalization-related competence protein ComEC/Rec2
ORF00888	SAG0783	269	hydrolase, haloacid dehalogenase-like family
ORF00889	SAG0784	314	sugar-binding transcriptional regulator, Lacl family
OBECCOOL	SAG0785	330	conserved hypothetical protein
ORF00890			
ORF00890 ORF00891	SAG0786	242	conserved domain protein

# Tabl 32: Conversion f ORF Ref N s. with SAG Ref Nos.

ORF Ref N .	SAGxxxx R f N .	aa	Ann tati n
ORF00893	SAG0788	202	superoxide dismutase, Fe-Mn
ORF00894	SAG0789	283	transcriptional antiterminator LicT
ORF00895	SAG0790	622	PTS system, beta-glucosides-specific IIABC components
ORF00896	SAG0791	475	6-phospho-beta-glucosidase
ORF00898	SAG0792	364	conserved hypothetical protein
ORF00899	SAG0793	380	conserved hypothetical protein TIGR00045
ORF00900	SAG0794	418	permease, GntP family
ORF00902	SAG0795	354	· · · · · · · · · · · · · · · · · · ·
ORF00903	SAG0796	147	conserved hypothetical protein transcriptional regulator, MarR family
ORF00904	SAG0797	342	S-adenosylmethionine:tRNA ribosyltransferase-
			isomerase
ORF00905	SAG0798	226	membrane protein, putative
ORF00906	SAG0799	233	glucosamine-6-phosphate isomerase
ORF00907	SAG0800	318	Glutathione S-transferases domain protein
ORF00908	SAG0801	239	ribosomal small subunit pseudouridine synthase
ORF00909	SAG0802	38	hypothetical protein
ORF00910	SAG0803	383	major facilitator family protein
ORF00911	SAG0804	315	competence protein CoiA
ORF00912	SAG0805	601	oligoendopeptidase B
ORF00913	SAG0806	208	hydrolase, haloacid dehalogenase-like family
ORF00914	SAG0807	235	O-methyltransferase family protein
ORF00916	· SAG0808	309	protease maturation protein, putative
ORF00918	SAG0809	161	conserved hypothetical protein
ORF00919	SAG0810	872	alanyl-tRNA synthetase
ORF00921	SAG0811	238	membrane protein, putative
ORF00922	SAG0812	272	glycosyl transferase, family 8
ORF00923	SAG0813	81	hypothetical protein
ORF00924	SAG0814	95	conserved domain protein
ORF00925	SAG0815	71	transcriptional regulator, Cro/Cl family
ORF00926	SAG0816	253	conserved hypothetical protein
ORF00927	SAG0817	187	conserved hypothetical protein
ORF00928	\$AG0818	319	ribonucleoside-diphosphate reductase 2, beta subunit
ORF00929	SAG0819	719	ribonucleoside-diphosphate reductase 2, alpha subunit
ORF00930	SAG0820	74	ribonucleoside-diphosphate reductase 2, NrdH-redoxin
ORF00931	SAG0821	87	phosphocarrier protein HPr
ORF00932	SAG0822	577	phosphoenolpyruvate-protein phosphotransferase
			, , , , , , , , , , , , , , , , , , , ,
ORF00933	SAG0823	475	glyceraldehyde-3-phosphate dehydrogenase, NADP-dependent
ORF00934	SAG0824	417	polysaccharide deacetylase family protein
ORF00935	SAG0825	360	ATP-dependent RNA helicase, DEAD/DEAH box family
ORF00936	SAG0826	209	uridine kinase
ORF00937	SAG0827	165	conserved hypothetical protein
ORF00938	SAG0828	554	DNA polymerase III, gamma and tau subunits
ORF00939	SAG0829	64	conserved hypothetical protein
ORF00940	SAG0830	311	biotin-acetyl-CoA-carboxylase ligas
ORF00941	SAG0831	398	S-adenosylmethlonine synthetase
ORF00942	SAG0832	753	hypothetical protein
ORF00943	SAG0833	181	hypothetical protein
ORF00944	SAG0834	42	hypothetical protein
ORF00945	SAG0835	188	conserved hypothetical protein



### Table 32: C nv rsion of ORF R f Nos. with SAG R f Nos.

ORF Ref N .	SAGxxxx Ref No.	aa	Annotation
ORF00946	SAG0836	184	conserved hypothetical protein
ORF00948	SAG0837	428	ABC transporter, ATP-binding protein
ORF00950	SAG0838	233	hypothetical protein
ORF00951	SAG0839	226	transcriptional regulator, TenA family
ORF00952	SAG0840	265	phosphomethylpyrimidine kinase
ORF00953 ,	SAG0841	256	hydroxyethylthiazole kinase
ORF00954	SAG0842	223	thiamine-phosphate pyrophosphorylase
ORF00955	SAG0843	419	UDP-N-acetylglucosamine 1-carboxyvinyltransferase
ORF00956	SAG0844	184	acetyltransferase, GNAT family
ORF00957	SAG0845	427	CBS domain protein
ORF00958	SAG0846	286	methionine aminopeptidase, type I
ORF00959	SAG0847	306	ribonuclease BN, putative
ORF00961	SAG0848	151	GtrA family protein
ORF00962	SAG0849	169	conserved hypothetical protein
ORF00963	SAG0850	652	DNA ligase, NAD-dependent
ORF00964	SAG0851	339	BmrU protein, putative
ORF00966	SAG0852	766	pullulanase, putative
ORF00967	SAG0853	622	1,4-alpha-glucan branching enzyme
ORF00968	SAG0854	379	glucose-1-phosphate adenylyltransferase
ORF00969	SAG0855		glycogen biosynthesis protein GlgD FRAMESHIFT
ORF00971	SAG0856	476	glycogen synthase ·
ORF00972	SAG0857	66	ATP synthase F0, C subunit
ORF00973	SAG0858	238	ATP synthase F0, A subunit
ORF00974	SAG0859	165	ATP synthase F0, B subunit
ORF00975	SAG0860	178	ATP synthase F1, delta subunit
ORF00976	SAG0861	501	ATP synthase F1, alpha subunit
ORF00977	SAG0862	293	ATP synthase F1, gamma subunit
ORF00978	SAG0863	468	ATP synthase F1, beta subunit
ORF00979	SAG0864	137	ATP synthase F1, epsilon subunit
ORF00980	SAG0865	76	conserved hypothetical protein
ORF00981	SAG0866	423	UDP-N-acetylglucosamine 1-carboxyvinyltransferase
ORF00982	SAG0867	63	conserved hypothetical protein
ORF00983	SAG0868	285	DNA-entry nuclease
ORF00984	SAG0869	346	phenylalanyl-tRNA synthetase, alpha subunit
ORF00985	SAG0870	173	acetyltransferase, GNAT family
ORF00986	SAG0871	801	phenylalanyl-tRNA synthetase, beta subunit
ORF00987	SAG0872	300	conserved hypothetical protein
ORF00988	SAG0873	1077	exonuclease RexB
ORF00989	SAG0874	1207	exonuclease RexA
ORF00990	SAG0875	305	magnesium transporter, CorA family, putative
ORF00991	SAG0876	458	tRNA modification GTPase TrmE
ORF00992	SAG0877	636	ABC transporter, ATP-binding protein
ORF00993	SAG0878	322	acetoin dehydrogenase, thymine PPi dependent, E1 component, alpha subunit
ORF00994	SAG0879	332	acetoin dehydrogenase, thymine PPi dependent, E1 component, beta subunit
ORF00995	SAG0880	462	acetoin dehydrogenase, thymine PPi dependent, E2 component, dihydrolipoamide acetyltransferase
ORF00996	SAG0881	585	acetoin dehydrogenase, thymine PPi dependent, E3 component, dihydrolipoamide dehydrogenase
ORF00997	SAG0882	329	lipoate-protein ligase A
ORF00998	SAG0883	261	cobyric acid synthase, putative



ORF Ref N .	SAGxxxx Ref No.	aa	Annotation
ORF00999	SAG0884	447	mur ligase family protein
ORF01000	SAG0885	283	conserved hypothetical protein TIGR00159
ORF01001	SAG0886	319	Gram-positive signal peptide, YSIRK family domain protein
ORF01002	SAG0887	450	phosphoglucomutase/phosphomannomutase family protein
ORF01003	SAG0888	123	conserved hypothetical protein
ORF01004	SAG0889	126	conserved hypothetical protein
ORF01005	SAG0890	376	oxygen-independent coproporphyrinogen III oxidase, putative
ORF01006	SAG0891	245	conserved hypothetical protein
ORF01007	SAG0892	256	hydrolase, haloacid dehalogenase-like family
ORF01008	SAG0893	218	conserved hypothetical protein
ORF01009	SAG0894	1370	conserved hypothetical protein
ORF01010	SAG0895	289	lipoyl-binding domain protein
ORF01011	SAG0896	108	oxidoreductase, putative
ORF01012	SAG0897	221	conserved hypothetical protein
ORF01013	SAG0898	83	hypothetical protein
ORF01014	SAG0899	57	hypothetical protein
ORF01015	SAG0900	56	hypothetical protein
ORF01016	SAG0901	127	hypothetical protein
ORF01018	SAG0902	45	hypothetical protein
ORF01019	SAG0903	44	hypothetical protein
ORF01021	SAG0904	56	hypothetical protein
ORF01022	SAG0905	138	nucleoside diphosphate kinase
ORF01023	SAG0906	610	GTP-binding protein LepA
ORF01024	SAG0907	877	streptococcal histidine triad family protein
ORF01025	SAG0908	203	HD domain protein
ORF01026	SAG0909	154	acetyltransferase, GNAT family
ORF01027	SAG0910	144	PilB-related protein
ORF01030	SAG0911	930	cation-transporting ATPase, E1-E2 family
ORF01031	SAG0912	367	nucleoside diphosphate kinase domain protein
ORF01032	SAG0913	212	chloramphenicol acetyltransferase
ORF01033	SAG0914	203	conserved hypothetical protein
ORF01034	SAG0915	405	Tn916, transposase
ORF01035	SAG0916	67	Tn916, excisionase
ORF01037	SAG0918	76	Tn916, hypothetical protein
ORF01038	SAG0919	157	Tn916, hypothetical protein
ORF01039	SAG0921	117	Tn916, transcriptional regulator, putative
ORF01040	SAG0923	639	Tn916, tetracycline resistance protein
ORF01041	SAG0925	310	Tn916, hypothetical protein
ORF01042	SAG0926	333	Tn916, NLP/P60 family protein
ORF01044	SAG0927	725	Tn916, hypothetical protein FRAMESHIFT
ORF01047	SAG0928	<del> </del>	Tn916, hypothetical protein FRAMESHIFT
ORF01048	SAG0929	168	Tn916, hypothetical protein
ORF01049	SAG0930	165	Tn916, hypothetical protein
ORF01050	SAG0931	73	Tn916, hypothetical protein
ORF01051	SAG0932	401	Tn916, transcriptional regulator, putative
ORF01052	SAG0933	461	Tn916, FtsK/SpollIE family protein
ORF01053	SAG0934	128	Tn916, hypothetical protein
ORF01054	SAG0935	104	Tn916, hypothetical protein
ORF01056	SAG0937		ABC transporter, ATP-binding protein, FRAMESHIFT
ORF01057	SAG0938	122	transcriptional regulator, GntR family
ORF01058	SAG0939	1034	DNA polymerase III, alpha subunit



### Tabl 32: Conv rsi n of ORF Ref Nos. with SAG Ref Nos.

ORFR fN.	SAGXXXX R f No.	aa	Ann tati n
ORF01059	SAG0940	340	6-phosphofructokinase
ORF01060	SAG0941	500	pyruvate kinase
ORF01061	SAG0942	185	signal peptidase I, putative
ORF01062	SAG0943	47	hypothetical protein
ORF01063	SAG0944	604	glucosaminefructose-6-phosphate aminotransferase
			(isomerizing)
ORF01064	SAG0945	377	IS1548, transposase
ORF01066	SAG0946	109	phnA protein
ORF01068	SAG0947	213	amino acid ABC transporter, permease protein
ORF01069	SAG0948	209	amino acid ABC transporter, ATP-binding protein
ORF01070	SAG0949	276	amino acid ABC transporter, amino acid-binding
			protein
ORF01072	SAG0950	82	ribosomal protein S20
ORF01073	SAG0951	306	pantothenate kinase
ORF01074	SAG0952	196	conserved hypothetical protein
ORF01075	SAG0953	129	cytidine deaminase
ORF01076	SAG0954	349	lipoprotein
ORF01077	SAG0955	511	sugar ABC transporter, ATP-binding protein
ORF01078	SAG0956	353	sugar ABC transporter, ATP-binding protein
01.11.07.0	OA00930	333	sugar ABC transporter, permease protein, putative
ORF01079	SAG0957	318	sugar ABC transporter, permease protein, putative
	0,10000.	0.0	sugar Abo transporter, permease protein, putative
ORF01080	SAG0958	456	NADH oxidase
ORF01081	SAG0959	329	L-lactate dehydrogenase
ORF01082	SAG0960	819	DNA gyrase, A subunit
ORF01083	SAG0961	247	sortase SrtA
ORF01084	SAG0962	137	glyoxylase family protein
ORF01085	SAG0963	320	conserved hypothetical protein
ORF01086	SAG0964	375	Na+/H+ exchanger family protein
ORF01087	SAG0965	127	IS1381, transposase OrfA
ORF01088	SAG0966	129	
ORF01089	SAG0967	520	IS1381, transposase OrfB GMP synthase
ORF01090	SAG0968	232	
ORF01091	SAG0969		transcriptional regulator, GntR family
ORF01091		444	gid protein
ORF01092	SAG0970	247	acetyltransferase, GNAT family
ORF01095	SAG0971	282	lipoprotein, putative
	SAG0972		conserved hypothetical protein, FRAMESHIFT
ORF01096	SAG0973	320	nisin-resistance protein, putative
ORF01097	. SAG0974	250	ABC transporter, ATP-binding protein
ORF01098	SAG0975	651	ABC transporter, permease protein, putative
ORF01099	SAG0976	222	DNA-binding response regulator
ORF01100	SAG0977	312	sensor histidine kinase
ORF01101	SAG0978	356	site-specific recombinase, phage integrase family
00504400			
ORF01102	SAG0979	553	ABC transporter, substrate binding protein, putative
00504400			
ORF01103	SAG0980	257	conserved hypothetical protein
ORF01104	SAG0981	228	SatD
ORF01106	SAG0982	521	signal recognition particle protein
ORF01108	SAG0983	110	conserved hypothetical protein
ORF01109	SAG0984	437	sensor histidine kinase CiaH
ORF01110	SAG0985	226	DNA-binding response regulator CiaR
ORF01111	SAG0986	849	aminopeptidase N
ORF01112	SAG0987	217	phosphate transport system regulatory protein PhoU

Table 32: Conversion of ORF Ref N s. with SAG R f Nos.

ORF Ref No.	SAGxxxx R fN.	aa	Ann tation
ORF01113	SAG0988	252	phosphate ABC transporter, ATP-binding protein PstB,
			putative
ORF01114	SAG0989	267	phosphate ABC transporter, ATP-binding protein PstB, putative
ORF01115	SAG0990	295	phosphate ABC transporter, permease protein PstA,
			putative
ORF01116	SAG0991	305	phosphate ABC transporter, permease protein
ORF01117	SAG0992	286	phosphate ABC transporter, phosphate-binding protein
ORF01118	SAG0993	436	NOL1/NOP2/sun family protein
ORF01119	SAG0994	254	inositol monophosphatase family protein
ORF01120	SAG0995	93	conserved hypothetical protein
ORF01121	SAG0996	137	conserved hypothetical protein
ORF01122	SAG0997	310	macrolide-efflux protein mreA/riboflavin biosynthesis protein RibF
ORF01123	SAG0998	294	tRNA pseudouridine synthase B
ORF01124	SAG0999	143	acetyltransferase, GNAT family
ORF01125	SAG1000	423	conserved hypothetical protein
ORF01126	SAG1001	196	conserved hypothetical protein
ORF01127	SAG1002	292	protease, putative
ORF01128	SAG1003	876	permease, putative
ORF01129	SAG1004	233	ABC transporter, ATP-binding protein
ORF01131	SAG1005	706	DNA topoisomerase I
ORF01132	SAG1006	280	DprA/SMF protein, putative DNA processing factor
ORF01133	SAG1007	342	iron-compound ABC transporter, iron-compound-
		0.2	binding protein
ORF01134	SAG1008	253	iron compound ABC transporter, ATP-binding protein
ORF01135	SAG1009	324	iron compound ABC transporter, permease protein
ORF01136	SAG1010	320	iron compound ABC transporter, permease protein
ORF01137	SAG1011	182	acetyltransferase, CysE/LacA/LpxA/NodL family
ORF01138	SAG1012	253	ribonuclease HII
ORF01139	SAG1013	283	GTP-binding protein
ORF01140	SAG1014	190	conserved hypothetical protein
ORF01142	SAG1015	494	carbon starvation protein CstA, putative
ORF01143	SAG1016	244	response regulator
ORF01144	SAG1017	579	sensor histidine kinase, putative
ORF01145	SAG1018	40	hypothetical protein
ORF01146	SAG1019	39	conserved hypothetical protein, FRAMESHIFT
ORF01148	SAG1020	227	hypothetical protein
ORF01149	SAG1021	107	hypothetical protein
ORF01150	SAG1022	177	hypothetical protein
ORF01151	SAG1023	48	hypothetical protein
ORF01152	SAG1024	183	hypothetical protein
ORF01153	SAG1025	149	hypothetical protein
ORF01156	SAG1026	1.70	immunogenic secreted protein, DEGENERATE
ORF01157	SAG1027	84	conserved hypothetical protein
ORF01158	SAG1028	196	hypothetical protein
ORF01159	SAG1028	101	hypothetical protein
ORF01160	SAG1029	304	conserved hypothetical protein
ORF01161	SAG1030	120	extracellular protein, putative POINT MUATION
ORF01162	SAG1031	85	
ORF01164	SAG1032 SAG1033		conserved hypothetical protein
ORF01166	SAG1033 SAG1034	1309	FtsK/SpollIE family protein
2.7.01100	SAG1034	55	hypothetical prot in



### Table 32: Conv rsion f ORF Ref N s. with SAG R f Nos.

ORF Ref N .	SAGxxxx R f No.	aa	Annotation
ORF01167	SAG1035	424	conserved hypothetical protein
ORF01168	SAG1036	80	conserved hypothetical protein
ORF01169	SAG1037	157	hypothetical protein
ORF01172	SAG1038	1003	phage infection protein, putative
ORF01173	SAG1039	96	conserved hypothetical protein
ORF01174	SAG1040	260	conserved domain protein
ORF01175	SAG1041	107	hypothetical protein
ORF01176	SAG1042	1060	
ORF01177	SAG1043	358	carbamoyl-phosphate synthase, large subunit
ORF01178	SAG1044	307	carbamoyl-phosphate synthase, small subunit
ORF01179	SAG1045		aspartate carbamoyltransferase
ORF01180	<del> </del>	430	dihydroorotase, multifunctional complex type
ORF01181	SAG1046 SAG1047	209	orotate phosphoribosyltransferase
ORF01182		233	orotidine 5'-phosphate decarboxylase
	SAG1048	410	membrane protein, putative
ORF01183	SAG1049	513	ABC transporter, ATP-binding protein
ORF01184	SAG1050	112	ribonucleotide reductase, truncation
ORF01185	SAG1051	358	aspartate-semialdehyde dehydrogenase
ORF01186	SAG1052	47	cell wall surface anchor family protein, putative
ORF01187	SAG1053	30	hypothetical protein
ORF01188	SAG1054	531	cardiolipin synthetase
ORF01189	SAG1055	556	formate-tetrahydrofolate ligase
ORF01190	SAG1056	339	lipoate-protein ligase A
ORF01191	SAG1057	292	conserved hypothetical protein
ORF01192	SAG1058	272	conserved hypothetical protein
ORF01193	SAG1059	110	glycine cleavage system H protein, putative
ORF01194	SAG1060	328	bacterial luciferase family protein
ORF01195	SAG1061	399	oxidoreductase, FMN-binding
ORF01197	SAG1062	282	lipoate-protein ligase A family protein
ORF01198	SAG1063	228	flavoprotein-related protein
ORF01199	SAG1064	180	flavoprotein family protein
ORF01200	SAG1065	190	membrane protein, putative
ORF01201	SAG1066	572	phosphoglucomutase
ORF01202	SAG1067	178	IS861, transposase OrfA
ORF01203	SAG1068	277	IS861, transposase OrfB
ORF01204	SAG1069	65	hypothetical protein
ORF01205	SAG1070	577	ABC transporter, ATP-binding/permease protein
ORF01206	SAG1071	573	ABC transporter, ATP-binding/permease protein
ORF01207	SAG1072	200	conserved hypothetical protein
ORF01208	SAG1073	325	conserved hypothetical protein
ORF01209	SAG1074	418	Serine hydroxymethyltransferase
ORF01210	SAG1075	183	Sua5/YciO/YrdC/YwlC family protein
ORF01211	SAG1076	276	modification methylase, HemK family
ORF01212	SAG1077	359	peptide chain release factor 1
ORF01213	SAG1078	189	thymidine kinases
ORF01214	SAG1079	60	4-oxalocrotonate tautomerase
ORF01215	SAG1080	47	hypothetical protein
ORF01216	SAG1081	312	ApbE family protein
ORF01217	SAG1082	200	conserved hypothetical protein
ORF01218	SAG1083	411	conserved hypothetical protein
ORF01219	SAG1084	262	formate/nitrite transporter family protein
ORF01220	SAG1085	424	xanthine permeas
ORF01221	SAG1086	193	xanthine phosphoribosyltransferase
ORF01222	SAG1087	327	guanosine monophosphate reductase
ORF01223	SAG1088	446	drug resistance transporter, EmrB/QacA family,
		1	putative



# Table 32: Conversion of ORF Ref Nos. with SAG R f Nos.

ORF Ref No.	SAGxxxx Ref N .	aa	Annotation
ORF01224	SAG1089	230	conserved hypothetical protein
ORF01225	SAG1090	666	potassium uptake protein, putative
ORF01226	SAG1091	216	oxidoreductase, short chain dehydrogenase/reductase family, FRAMESHIFT
ORF01227	SAG1092	330	phosphate acetyltransferase
ORF01228	SAG1093	294	ribosomal large subunit pseudouridine synthase, RluD subfamily
ORF01229	SAG1094	278	conserved hypothetical protein
ORF01230	SAG1095	223	GTP pyrophosphokinase family protein
ORF01231	SAG1096	190	conserved hypothetical protein
ORF01232	SAG1097	324	ribose-phosphate pyrophosphokinase
ORF01233	SAG1098	371	cysteine desulphurase
ORF01234	SAG1099	115	conserved hypothetical protein
ORF01235	SAG1100	210	DNA-binding protein
ORF01236	SAG1101	226	DNA repair protein RadC
ORF01237	SAG1102	377	membrane protein, putative
ORF01238	SAG1103	478	6-phospho-beta-glucosidase
ORF01239	SAG1104	204	platelet activating factor, putative
ORF01240	SAG1105	273	hydrolase, haloacid dehalogenase-like family
ORF01241	SAG1106	309	transcriptional regulator, AraC family, putative
ORF01242	SAG1107	510	voltage-gated chloride channel family protein
ORF01243	SAG1108	357	spermidine/putrescine ABC transporter,
			spermidine/putrescine-binding protein
ORF01244	SAG1109	258	spermidine/putrescine ABC transporter, permease protein
ORF01245	SAG1110	264	spermidine/putrescine ABC transporter, permease protein
ORF01246	SAG1111	384	spermidine/putrescine ABC transporter, ATP-binding protein
ORF01247	SAG1112	300	UDP-N-acetylenolpyruvoylglucosamine reductase
ORF01248	SAG1113	162	2-amino-4-hydroxy-6-hydroxymethyldihydropteridine pyrophosphokinase
ORF01249	SAG1114	120	dihydroneopterin aldolase
ORF01250	SAG1115	267	dihydropteroate synthase
ORF01251	SAG1116	187	GTP cyclohydrolase I
ORF01252	SAG1117	420	folylpolyglutamate synthase
ORF01253	SAG1118	295	rarD protein
ORF01254	SAG1119	288	homoserine kinase
ORF01255	SAG1120	427	homoserine dehydrogenase
ORF01256	SAG1121	295	polysaccharide deacetylase family protein
ORF01257	SAG1122	515	transporter, BCCT family protein
ORF01258	SAG1123	34	hypothetical protein
ORF01259	SAG1124	458	aldehyde dehydrogenase family protein
ORF01260	SAG1125	335	membrane protein
ORF01261	SAG1126	228	conserved hypothetical protein
ORF01262	SAG1127	113	conserved hypothetical protein, FRAMESHIFT
ORF01263		187	hypothetical protein
ORF01264	SAG1128	65	transcriptional regulator, Cro/Cl family
	SAG1129	36	hypothetical protein
ORF01265			Tarana bratani
ORF01265 ORF01266	SAG1130	49	hypothetical protein
		49 164	hypothetical protein thiol peroxidase
ORF01266	SAG1130 SAG1131 SAG1132	49 164 219	hypothetical protein thiol peroxidase conserved hypothetical protein



ORF01316 SAG1175 485 capsular polysaccharide biosynthesis protein CpsA ORF01317 SAG1176 290 capsular polysaccharide synthesis operon transcriptional regulator CpsY ORF01318 SAG1177 255 cpsIaS protein ORF01319 SAG1178 236 purine nucleoside phosphorylase	ORF Ref No.	SAGxxxx R f No.	aa	Ann tation
Uptake protein, TrkA family	ORF01273	SAG1134	213	transcriptional regulator, GntR family/potassioum
ORF01274         SAG1135         183         gls24 protein, putative           ORF01276         SAG1137         180         gls24 protein, putative           ORF01277         SAG1138         64         conserved hypothetical protein           ORF01277         SAG1138         64         conserved hypothetical protein           ORF01280         SAG1140         82         conserved hypothetical protein           ORF01281         SAG1141         112         conserved hypothetical protein           ORF01282         SAG1142         759         ATP-dependent DNA heticase PcrA           ORF01283         SAG1143         100         conserved hypothetical protein, FRAMESHIFT           ORF01284         SAG1144         441         uracil permease           ORF01285         SAG1144         441         uracil permease           ORF01286         SAG1148         441         cation efflux family protein           ORF01287         SAG1148         231         membrane protein, putative           ORF01288         SAG1148         231         membrane protein, putative           ORF01289         SAG1150         400         ribosomal protein S1           ORF01290         SAG1151         400         ribosomal protein           ORF01				uptake protein, TrkA family
ORF01276         SAG1136         conserved hypothetical protein FRAMESHIFT           ORF01277         SAG1137         180         gis24 protein, putative           ORF01279         SAG1138         64         conserved hypothetical protein           ORF01279         SAG1139         193         conserved hypothetical protein           ORF01280         SAG1140         82         conserved hypothetical protein           ORF01281         SAG1141         112         conserved hypothetical protein           ORF01282         SAG1141         172         conserved hypothetical protein           ORF01283         SAG1141         112         conserved hypothetical protein, FRAMESHIFT           ORF01284         SAG1144         441         uracil permease           ORF01285         SAG1144         441         uracil permease           ORF01286         SAG1144         441         uracil permease           ORF01287         SAG1147         130         conserved hypothetical protein           ORF01288         SAG1148         231         membrane protein, putative           ORF01289         SAG1150         400         ribosomal protein S1           ORF01290         SAG1151         76         conserved hypothetical protein           ORF01	ORF01274	SAG1135	183	
ORF01276         SAG1137         180         gls24 protein, putative           ORF01277         SAG1138         64         conserved hypothetical protein           ORF01280         SAG1140         82         conserved hypothetical protein           ORF01281         SAG1141         112         conserved hypothetical protein           ORF01282         SAG1142         759         ATP-dependent DNA helicase PcrA           ORF01283         SAG1143         100         conserved hypothetical protein, FRAMESHIFT           ORF01284         SAG1144         441         uracil permease           ORF01285         SAG1144         441         uracil permease           ORF01286         SAG1147         130         conserved hypothetical protein           ORF01287         SAG1144         411         cation efflux family protein           ORF01288         SAG1148         231         membrane protein, putative           ORF01289         SAG1149         207         conserved hypothetical protein           ORF01291         SAG1150         440         membrane protein, putative           ORF01292         SAG1151         76         conserved hypothetical protein           ORF01293         SAG1152         340         branched-chain amino acid aminotransferase	ORF01275	SAG1136		
ORF01277         SAG1138         64         conserved hypothetical protein           ORF01280         SAG1140         82         conserved hypothetical protein           ORF01281         SAG1141         112         conserved hypothetical protein           ORF01282         SAG1141         112         conserved hypothetical protein           ORF01283         SAG1143         100         conserved hypothetical protein, FRAMESHIFT           ORF01284         SAG1144         441         uracil permease           ORF01285         SAG1144         441         uracil permease           ORF01286         SAG1146         441         cation efflux family protein           ORF01287         SAG1147         130         conserved hypothetical protein           ORF01288         SAG1148         231         membrane protein, putative           ORF01289         SAG1149         207         conserved hypothetical protein           ORF01290         SAG1150         400         ribosomal protein S1           ORF01291         SAG1151         76         conserved hypothetical protein           ORF01292         SAG1153         819         DNA topoisomerase IV, A subunit           ORF01293         SAG1155         207         conserved hypothetical protein     <	ORF01276	SAG1137	180	
ORF01279         SAG1139         193         conserved hypothetical protein           ORF01280         SAG1140         82         conserved hypothetical protein           ORF01281         SAG1141         112         conserved hypothetical protein           ORF01282         SAG1142         759         ATP-dependent DNA helicase PcrA           ORF01283         SAG1143         100         conserved hypothetical protein, FRAMESHIFT           ORF01284         SAG1144         441         uracil permease           ORF01285         SAG1146         441         calium allanine symporter family protein           ORF01286         SAG1148         441         calium efflux family protein           ORF01287         SAG1148         231         membrane protein, putative           ORF01288         SAG1148         231         membrane protein, putative           ORF01289         SAG1148         231         membrane protein, putative           ORF01280         SAG1150         400         ribosomal protein s1           ORF01291         SAG1151         76         conserved hypothetical protein           ORF01292         SAG1153         340         branched-chain amino acid aminotransferase           ORF01293         SAG1155         207         conserved h	ORF01277	SAG1138	64	
ORF01280         SAG1140         82         conserved hypothetical protein           ORF01281         SAG1141         112         conserved hypothetical protein           ORF01282         SAG1142         759         ATP-dependent DNA helicase PcrA           ORF01283         SAG1143         100         conserved hypothetical protein, FRAMESHIFT           ORF01284         SAG1144         441         uracii permease           ORF01285         SAG1145         448         sodiumtalanine symporter family protein           ORF01286         SAG1146         411         cation efflux family protein           ORF01287         SAG1147         130         conserved hypothetical protein           ORF01289         SAG1148         231         membrane protein, putative           ORF01289         SAG1149         207         conserved hypothetical protein           ORF01291         SAG1150         400         ribosomal protein S1           ORF01292         SAG1151         76         conserved hypothetical protein           ORF01293         SAG1152         340         branched-chain amino acid aminotransferase           ORF01294         SAG1153         819         DNA topoisomerase IV, A subunit           ORF01295         SAG1154         653         DNA	ORF01279			
ORF01281         SAG1141         112         conserved hypothetical protein           ORF01282         SAG1142         759         ATP-dependent DNA helicase PcrA           ORF01283         SAG1143         100         conserved hypothetical protein, FRAMESHIFT           ORF01284         SAG1144         441         uracil permease           ORF01285         SAG1146         441         cation efflux family protein           ORF01286         SAG1148         441         cation efflux family protein           ORF01287         SAG1148         441         cation efflux family protein           ORF01288         SAG1148         231         membrane protein, putative           ORF01289         SAG1140         207         conserved hypothetical protein           ORF01290         SAG1150         400         ribosomal protein, putative           ORF01291         SAG1150         400         ribosomal protein, putative           ORF01292         SAG1151         76         conserved hypothetical protein           ORF01293         SAG1152         340         branched-chain amino acid aminotransferase           ORF01294         SAG1153         319         DNA topolsomerase IV, a subunit           ORF01295         SAG1154         653         DNA topolso				
ORF01282         SAG1142         759         ATP-dependent DNA helicase PcrA           ORF01283         SAG1143         100         conserved hypothetical protein, FRAMESHIFT           ORF01285         SAG1144         441         uracil permease           ORF01286         SAG1145         448         sodium:alanine symporter family protein           ORF01287         SAG1147         130         conserved hypothetical protein           ORF01288         SAG1148         231         membrane protein, putative           ORF01289         SAG1149         207         conserved hypothetical protein           ORF01289         SAG1150         400         ribosomal protein S1           ORF01291         SAG1151         76         conserved hypothetical protein           ORF01292         SAG1153         819         DNA topoisomerase IV, A subunit           ORF01293         SAG1153         819         DNA topoisomerase IV, B subunit           ORF01294         SAG1153         819         DNA topoisomerase IV, B subunit           ORF01295         SAG1155         207         conserved hypothetical protein TIGR00023           ORF01296         SAG1155         217         uracil-DNA glycosylase           ORF01297         SAG1166         217         uracil-D	ORF01281			
ORF01283         SAG1143         100         conserved hypothetical protein, FRAMESHIFT           ORF01284         SAG1144         441         uracil permease           ORF01285         SAG1145         448         sodium:alanine symporter family protein           ORF01286         SAG1146         441         cation efflux family protein           ORF01287         SAG1147         130         conserved hypothetical protein           ORF01288         SAG1148         231         membrane protein, putative           ORF01289         SAG1149         207         conserved hypothetical protein           ORF01290         SAG1150         400         ribosomal protein S1           ORF01291         SAG1151         76         conserved hypothetical protein           ORF01292         SAG1152         340         branched-chain amino acid aminotransferase           ORF01293         SAG1153         819         DNA topolsomerase IV, A subunit           ORF01294         SAG1153         819         DNA topolsomerase IV, B subunit           ORF01295         SAG1155         207         conserved hypothetical protein TIGR00023           ORF01297         SAG1156         217         uracil-DNA glycosylase           ORF01298         SAG1157         161	ORF01282			
ORF01284         SAG1144         441         uracil permease           ORF01285         SAG1145         448         sodium:alanine symporter family protein           ORF01286         SAG1145         448         sodium:alanine symporter family protein           ORF01287         SAG1147         130         conserved hypothetical protein           ORF01288         SAG1148         231         membrane protein, putative           ORF01289         SAG1149         207         conserved hypothetical protein           ORF01290         SAG1150         400         ribosomal protein S1           ORF01291         SAG1151         76         conserved hypothetical protein           ORF01292         SAG1152         340         branched-chain amino acid aminotransferase           ORF01294         SAG1153         819         DNA topolsomerase IV, A subunit           ORF01295         SAG1154         653         DNA topolsomerase IV, A subunit           ORF01296         SAG1155         207         conserved hypothetical protein TIGR00023           ORF01297         SAG1156         217         uracil-DNA glycosylase           ORF01298         SAG1157         161         conserved hypothetical protein           ORF01299         SAG1158         413         C				
ORF01285         SAG1145         448         sodium:alianine symporter family protein           ORF01286         SAG1146         411         cation efflux family protein           ORF01287         SAG1147         130         conserved hypothetical protein           ORF01288         SAG1148         231         membrane protein, putative           ORF01289         SAG1149         207         conserved hypothetical protein           ORF01290         SAG1150         400         ribosomal protein S1           ORF01291         SAG1151         76         conserved hypothetical protein           ORF01292         SAG1151         76         conserved hypothetical protein           ORF01293         SAG1153         340         branched-chain amino acid aminotransferase           ORF01294         SAG1153         819         DNA topolsomerase IV, A subunit           ORF01295         SAG1154         653         DNA topolsomerase IV, A subunit           ORF01296         SAG1155         207         conserved hypothetical protein TIGR00023           ORF01297         SAG1166         217         uracii-DNA glycosylase           ORF01298         SAG1158         413         CMP-N-acetylneuraminic acid synthetase NeuA           ORF01299         SAG1158         413 </td <td></td> <td></td> <td></td> <td></td>				
ORF01286         SAG1146         411         cation efflux family protein           ORF01287         SAG1147         130         conserved hypothetical protein           ORF01288         SAG1148         231         membrane protein, putative           ORF01289         SAG1149         207         conserved hypothetical protein           ORF01290         SAG1150         400         ribosomal protein S1           ORF01291         SAG1151         76         conserved hypothetical protein           ORF01292         SAG1151         76         conserved hypothetical protein           ORF01293         SAG1151         76         conserved hypothetical protein           ORF01294         SAG1153         819         DNA topolsomerase IV, A subunit           ORF01295         SAG1154         653         DNA topolsomerase IV, B subunit           ORF01296         SAG1155         207         conserved hypothetical protein TIGR00023           ORF01297         SAG1156         217         conserved hypothetical protein TIGR00023           ORF01298         SAG1157         161         conserved hypothetical protein TIGR00023           ORF01299         SAG1158         413         CMP-N-acetylineuraminic acid synthetase NeuA           ORF01300         SAG1160 <td< td=""><td></td><td></td><td></td><td></td></td<>				
ORF01287         SAG1147         130         conserved hypothetical protein           ORF01288         SAG1148         231         membrane protein, putative           ORF01289         SAG1149         207         conserved hypothetical protein           ORF01290         SAG1150         400         ribosomal protein S1           ORF01291         SAG1151         76         conserved hypothetical protein           ORF01292         SAG1152         340         branched-chain amino acid aminotransferase           ORF01294         SAG1153         819         DNA topoisomerase IV, A subunit           ORF01295         SAG1154         653         DNA topoisomerase IV, B subunit           ORF01296         SAG1155         207         conserved hypothetical protein TIGR00023           ORF01297         SAG1166         217         uracii-DNA glycosylase           ORF01298         SAG1157         161         conserved hypothetical protein           ORF01299         SAG1158         413         CMP-N-acetylaeuraminic acid synthetase NeuA           ORF01300         SAG1159         209         neuD protein           ORF01301         SAG1160         384         UDP-N-acetylglucosamine-2-epimerase NeuC           ORF01302         SAG1161         341 <td< td=""><td></td><td></td><td></td><td></td></td<>				
ORF01288         SAG1148         231         membrane protein, putative           ORF01289         SAG1149         207         conserved hypothetical protein           ORF01290         SAG1150         400         ribosomal protein S1           ORF01291         SAG1151         76         conserved hypothetical protein           ORF01292         SAG1152         340         branched-chain amino acid aminotransferase           ORF01294         SAG1153         819         DNA topolsomerase IV, A subunit           ORF01295         SAG1154         653         DNA topolsomerase IV, B subunit           ORF01296         SAG1155         207         conserved hypothetical protein TIGR00023           ORF01297         SAG1166         217         uracil-DNA glycosylase           ORF01298         SAG1157         161         conserved hypothetical protein TIGR00023           ORF01299         SAG1158         413         CMP-N-acetylneuraminic acid synthetase NeuA           ORF01300         SAG1159         209         neuD protein           ORF01301         SAG1160         384         UDP-N-acetylglucosamine-2-epimerase NeuC           ORF01302         SAG1161         341         N-acetyl neuramic acid synthetase NeuB           ORF01303         SAG1163         31				
ORF01289         SAG1149         207         conserved hypothetical protein           ORF01290         SAG1150         400         ribosomal protein S1           ORF01291         SAG1151         76         conserved hypothetical protein           ORF01292         SAG1151         76         conserved hypothetical protein           ORF01293         SAG1153         819         DNA topoisomerase IV, A subunit           ORF01295         SAG1154         653         DNA topoisomerase IV, B subunit           ORF01298         SAG1155         207         conserved hypothetical protein TIGR00023           ORF01297         SAG1166         217         uracil-DNA glycosylase           ORF01298         SAG1157         161         conserved hypothetical protein TIGR00023           ORF01299         SAG1158         413         CMP-N-acetylneuraminic acid synthetase NeuA           ORF01300         SAG1169         384         UDP-N-acetylglucosamine-2-epimerase NeuC           ORF01301         SAG1160         384         UDP-N-acetylglucosamine-2-epimerase NeuC           ORF01302         SAG1161         341         N-acetyl neuramic acid synthetase NeuB           ORF01303         SAG1162         466         cpsL protein           ORF01304         SAG1163         3				
ORF01290         SAG1150         400         ribosomal protein S1           ORF01291         SAG1151         76         conserved hypothetical protein           ORF01292         SAG1152         340         branched-chain amino acid aminotransferase           ORF01294         SAG1153         819         DNA topoisomerase IV, A subunit           ORF01295         SAG1154         653         DNA topoisomerase IV, B subunit           ORF01298         SAG1155         207         conserved hypothetical protein TIGR00023           ORF01297         SAG1166         217         uracil-DNA glycosylase           ORF01298         SAG1157         161         conserved hypothetical protein           ORF01299         SAG1158         413         CMP-N-acetylneuraminic acid synthetase NeuA           ORF01300         SAG1159         209         neuD protein           ORF01301         SAG1160         384         UDP-N-acetylglucosamine-2-epimerase NeuC           ORF01302         SAG1161         341         N-acetyl neuramic acid synthetase NeuB           ORF01303         SAG1162         466         cpsL protein           ORF01304         SAG1163         318         cpsVK protein           ORF01305         SAG1164         321         cpsVV protein				
ORF01291         SAG1151         76         conserved hypothetical protein           ORF01292         SAG1152         340         branched-chain amino acid aminotransferase           ORF01294         SAG1153         819         DNA topoisomerase IV, A subunit           ORF01295         SAG1154         653         DNA topoisomerase IV, A subunit           ORF01298         SAG1155         207         conserved hypothetical protein TIGR00023           ORF01297         SAG1156         217         uracil-DNA glycosylase           ORF01298         SAG1157         161         conserved hypothetical protein TIGR00023           ORF01299         SAG1158         413         CMP-N-acetylneuraminic acid synthetase NeuA           ORF01300         SAG1159         209         neuD protein           ORF01301         SAG1169         384         UDP-N-acetylglucosamine-2-epimerase NeuC           ORF01302         SAG1160         384         UDP-N-acetylglucosamine-2-epimerase NeuC           ORF01303         SAG1162         466         cpsL protein           ORF01304         SAG1163         318         cpsVK protein           ORF01305         SAG1164         321         cpsVK protein           ORF01306         SAG1166         295         cpsVN protein <td></td> <td></td> <td></td> <td></td>				
ORF01292         SAG1152         340         branched-chain amino acid aminotransferase           ORF01294         SAG1153         819         DNA topoisomerase IV, A subunit           ORF01295         SAG1154         653         DNA topoisomerase IV, B subunit           ORF01296         SAG1155         207         conserved hypothetical protein TIGR00023           ORF01297         SAG1156         217         uracil-DNA glycosylase           ORF01298         SAG1157         161         conserved hypothetical protein           ORF01299         SAG1158         413         CMP-N-acetylneuraminic acid synthetase NeuA           ORF01300         SAG1159         209         neuD protein           ORF01301         SAG1160         384         UDP-N-acetylglucosamine-2-epimerase NeuC           ORF01302         SAG1161         341         N-acetyl neuramic acid synthetase NeuB           ORF01303         SAG1161         341         N-acetyl neuramic acid synthetase NeuB           ORF01304         SAG1163         318         cpsVK protein           ORF01305         SAG1164         321         cpsVK protein           ORF01306         SAG1165         327         cpsVO protein           ORF01307         SAG1166         295         cpsVn protein				
ORF01294         SAG1153         819         DNA topoisomerase IV, A subunit           ORF01295         SAG1154         653         DNA topoisomerase IV, B subunit           ORF01298         SAG1155         207         conserved hypothetical protein TIGR00023           ORF01297         SAG1156         217         uracii-DNA glycosylase           ORF01298         SAG1157         161         conserved hypothetical protein           ORF01299         SAG1158         413         CMP-N-acetylneuraminic acid synthetase NeuA           ORF01300         SAG1159         209         neuD protein           ORF01301         SAG1160         384         UDP-N-acetylglucosamine-2-epimerase NeuC           ORF01302         SAG1161         341         N-acetyl neuramic acid synthetase NeuB           ORF01303         SAG1162         466         cpsL protein           ORF01304         SAG1163         318         cpsVK protein           ORF01305         SAG1164         321         cpsVJ protein           ORF01306         SAG1165         327         cpsVO protein           ORF01307         SAG1166         295         cpsVN protein           ORF01310         SAG1168         364         cpsVM protein           ORF01311 <t< td=""><td></td><td></td><td></td><td></td></t<>				
ORF01295         SAG1154         653         DNA topolsomerase IV, B subunit           ORF01296         SAG1155         207         conserved hypothetical protein TIGR00023           ORF01297         SAG1156         217         uracil-DNA glycosylase           ORF01298         SAG1157         161         conserved hypothetical protein           ORF01299         SAG1158         413         CMP-N-acetylneuraminic acid synthetase NeuA           ORF01300         SAG1159         209         neuD protein           ORF01301         SAG1160         384         UDP-N-acetylglucosamine-2-epimerase NeuC           ORF01302         SAG1161         341         N-acetyl neuramic acid synthetase NeuB           ORF01302         SAG1161         341         N-acetyl neuramic acid synthetase NeuB           ORF01302         SAG1161         341         N-acetyl neuramic acid synthetase NeuB           ORF01303         SAG1162         466         cpst protein           ORF01304         SAG1163         318         cpsVK protein           ORF01305         SAG1163         318         cpsVK protein           ORF01306         SAG1165         327         cpsVO protein           ORF01307         SAG1166         295         cpsVN protein				
ORF01296         SAG1155         207         conserved hypothetical protein TIGR00023           ORF01297         SAG1156         217         uracii-DNA glycosylase           ORF01298         SAG1157         161         conserved hypothetical protein           ORF01299         SAG1158         413         CMP-N-acetylneuraminic acid synthetase NeuA           ORF01300         SAG1159         209         neuD protein           ORF01301         SAG1160         384         UDP-N-acetylglucosamine-2-epimerase NeuC           ORF01302         SAG1161         341         N-acetyl neuramic acid synthetase NeuB           ORF01303         SAG1162         466         cps. protein           ORF01304         SAG1163         318         cpsVK protein           ORF01305         SAG1163         318         cpsVK protein           ORF01306         SAG1164         321         cpsVO protein           ORF01307         SAG1166         295         cpsVN protein           ORF01308         SAG1167         241         cpsVM protein           ORF01310         SAG1168         364         cpsVH protein           ORF01311         SAG1170         149         CpsE           ORF01312         SAG1171         462         <				
ORF01297         SAG1156         217         uracil-DNA glycosylase           ORF01298         SAG1157         161         conserved hypothetical protein           ORF01299         SAG1158         413         CMP-N-acetylneuraminic acid synthetase NeuA           ORF01300         SAG1159         209         neuD protein           ORF01301         SAG1160         384         UDP-N-acetylglucosamine-2-epimerase NeuC           ORF01302         SAG1161         341         N-acetyl neuramic acid synthetase NeuB           ORF01303         SAG1162         466         cpsL protein           ORF01304         SAG1163         318         cpsVK protein           ORF01305         SAG1164         321         cpsVJ protein           ORF01306         SAG1165         327         cpsVO protein           ORF01307         SAG1166         295         cpsVN protein           ORF01308         SAG1166         295         cpsVN protein           ORF01310         SAG1168         364         cpsVH protein           ORF01310         SAG1168         364         cpsVH protein           ORF01311         SAG1170         149         CpsF           ORF01312         SAG1171         462         CpsE				
ORF01298         SAG1157         161         conserved hypothetical protein           ORF01299         SAG1158         413         CMP-N-acetylneuraminic acid synthetase NeuA           ORF01300         SAG1169         209         neuD protein           ORF01301         SAG1160         384         UDP-N-acetylglucosamine-2-epimerase NeuC           ORF01302         SAG1161         341         N-acetyl neuramic acid synthetase NeuB           ORF01302         SAG1161         341         N-acetyl neuramic acid synthetase NeuB           ORF01303         SAG1161         341         N-acetyl neuramic acid synthetase NeuB           ORF01303         SAG1162         466         cps. protein           ORF01304         SAG1163         318         cps. Vy protein           ORF01305         SAG1164         321         cps. Vy protein           ORF01306         SAG1165         327         cps. Vy protein           ORF01307         SAG1166         295         cps. Vy protein           ORF01308         SAG1167         241         cps. Vy protein           ORF01310         SAG1169         163         Cps. Vg           ORF01311         SAG1170         149         Cps. Vg           ORF01312         SAG1171         <				
ORF01299         SAG1158         413         CMP-N-acetylneuraminic acid synthetase NeuA           ORF01300         SAG1159         209         neuD protein           ORF01301         SAG1160         384         UDP-N-acetylglucosamine-2-epimerase NeuC           ORF01302         SAG1161         341         N-acetyl neuramic acid synthetase NeuB           ORF01303         SAG1162         466         cps. protein           ORF01304         SAG1163         318         cpsVK protein           ORF01305         SAG1163         321         cpsVK protein           ORF01306         SAG1165         327         cpsVO protein           ORF01307         SAG1166         295         cpsVN protein           ORF01308         SAG1167         241         cpsVM protein           ORF01309         SAG1168         364         cpsVH protein           ORF01310         SAG1169         163         CpsVG           ORF01311         SAG1170         149         CpsF           ORF01312         SAG1171         462         CpsE           ORF01313         SAG1172         229         CpsD protein           ORF01316         SAG1175         485         capsular polysaccharide biosynthesis protein CpsA				
ORF01300         SAG1159         209         neuD protein           ORF01301         SAG1160         384         UDP-N-acetylglucosamine-2-epimerase NeuC           ORF01302         SAG1161         341         N-acetyl neuramic acid synthetase NeuB           ORF01303         SAG1162         466         cpsL protein           ORF01304         SAG1163         318         cpsVK protein           ORF01305         SAG1164         321         cpsVJ protein           ORF01306         SAG1165         327         cpsVO protein           ORF01307         SAG1166         295         cpsVN protein           ORF01308         SAG1167         241         cpsVM protein           ORF01309         SAG1168         364         cpsVM protein           ORF01310         SAG1169         163         CpsVG           ORF01311         SAG1170         149         CpsF           ORF01312         SAG1171         462         CpsE           ORF01314         SAG1173         230         cpsC protein           ORF01315         SAG1174         243         capsular polysaccharide biosynthesis protein CpsE           ORF01316         SAG1176         290         capsular polysaccharide synthesis operon transcriptional regul				
ORF01301         SAG1160         384         UDP-N-acetylglucosamine-2-epimerase NeuC           ORF01302         SAG1161         341         N-acetyl neuramic acid synthetase NeuB           ORF01303         SAG1162         466         cpsL protein           ORF01304         SAG1163         318         cpsVK protein           ORF01305         SAG1164         321         cpsVJ protein           ORF01306         SAG1165         327         cpsVO protein           ORF01307         SAG1166         295         cpsVN protein           ORF01308         SAG1166         295         cpsVN protein           ORF01309         SAG1168         364         cpsVH protein           ORF01310         SAG1169         163         CpsVG           ORF01311         SAG1170         149         CpsF           ORF01312         SAG1171         462         CpsE           ORF01313         SAG1172         229         CpsD protein           ORF01314         SAG1173         230         cpsC protein           ORF01315         SAG1174         243         capsular polysaccharide biosynthesis protein CpsE           ORF01316         SAG1176         290         capsular polysaccharide synthesis operon transcriptional regul				
ORF01302         SAG1161         341         N-acetyl neuramic acid synthetase NeuB           ORF01303         SAG1162         466         cpsL protein           ORF01304         SAG1163         318         cpsVK protein           ORF01305         SAG1164         321         cpsVJ protein           ORF01306         SAG1165         327         cpsVO protein           ORF01307         SAG1166         295         cpsVN protein           ORF01308         SAG1167         241         cpsVM protein           ORF01309         SAG1168         364         cpsVH protein           ORF01310         SAG1169         163         CpsVG           ORF01311         SAG1170         149         CpsF           ORF01312         SAG1171         462         CpsE           ORF01313         SAG1172         229         CpsD protein           ORF01314         SAG1173         230         cpsC protein           ORF01315         SAG1174         243         capsular polysaccharide biosynthesis protein CpsE           ORF01316         SAG1176         290         capsular polysaccharide synthesis operon transcriptional regulator CpsY           ORF01318         SAG1177         255         cpsIaS protein				
ORF01303         SAG1162         466         cpsL protein           ORF01304         SAG1163         318         cpsVK protein           ORF01305         SAG1164         321         cpsVJ protein           ORF01306         SAG1165         327         cpsVO protein           ORF01307         SAG1166         295         cpsVN protein           ORF01308         SAG1167         241         cpsVM protein           ORF01309         SAG1168         364         cpsVH protein           ORF01310         SAG1169         163         CpsVG           ORF01311         SAG1170         149         CpsF           ORF01312         SAG1171         462         CpsE           ORF01313         SAG1172         229         CpsD protein           ORF01314         SAG1173         230         cpsC protein           ORF01315         SAG1174         243         capsular polysaccharide biosynthesis protein CpsE           ORF01316         SAG1175         485         capsular polysaccharide synthesis operon transcriptional regulator CpsY           ORF01318         SAG1177         255         cpslaS protein           ORF01319         SAG1178         236         purine nucleoside phosphorylase		SAG1160	384	
ORF01304         SAG1163         318         cpsVK protein           ORF01305         SAG1164         321         cpsVJ protein           ORF01306         SAG1165         327         cpsVO protein           ORF01307         SAG1166         295         cpsVN protein           ORF01308         SAG1166         295         cpsVN protein           ORF01309         SAG1168         364         cpsVH protein           ORF01310         SAG1169         163         CpsVG           ORF01311         SAG1170         149         CpsF           ORF01312         SAG1171         462         CpsE           ORF01313         SAG1172         229         CpsD protein           ORF01314         SAG1173         230         cpsC protein           ORF01315         SAG1174         243         capsular polysaccharide biosynthesis protein CpsE           ORF01316         SAG1175         485         capsular polysaccharide synthesis operon transcriptional regulator CpsY           ORF01318         SAG1177         255         cpslaS protein           ORF01319         SAG1178         236         purine nucleoside phosphorylase			341	
ORF01305         SAG1164         321         cpsVJ protein           ORF01306         SAG1165         327         cpsVO protein           ORF01307         SAG1166         295         cpsVN protein           ORF01308         SAG1167         241         cpsVM protein           ORF01309         SAG1168         364         cpsVH protein           ORF01310         SAG1169         163         CpsVG           ORF01311         SAG1170         149         CpsF           ORF01312         SAG1171         462         CpsE           ORF01313         SAG1172         229         CpsD protein           ORF01314         SAG1173         230         cpsC protein           ORF01315         SAG1174         243         capsular polysaccharide biosynthesis protein CpsE           ORF01316         SAG1175         485         capsular polysaccharide biosynthesis operon transcriptional regulator CpsY           ORF01318         SAG1177         255         cpslaS protein           ORF01319         SAG1178         236         purine nucleoside phosphorylase			466	
ORF01306         SAG1165         327         cpsVO protein           ORF01307         SAG1166         295         cpsVN protein           ORF01308         SAG1167         241         cpsVM protein           ORF01309         SAG1168         364         cpsVH protein           ORF01310         SAG1169         163         CpsVG           ORF01311         SAG1170         149         CpsF           ORF01312         SAG1171         462         CpsE           ORF01313         SAG1172         229         CpsD protein           ORF01314         SAG1173         230         cpsC protein           ORF01315         SAG1174         243         capsular polysaccharide biosynthesis protein CpsE           ORF01316         SAG1175         485         capsular polysaccharide biosynthesis protein CpsE           ORF01317         SAG1176         290         capsular polysaccharide synthesis operon transcriptional regulator CpsY           ORF01318         SAG1177         255         cpslaS protein           ORF01319         SAG1178         236         purine nucleoside phosphorylase	ORF01304	SAG1163	318	cpsVK protein
ORF01307         SAG1166         295         cpsVN protein           ORF01308         SAG1167         241         cpsVM protein           ORF01309         SAG1168         364         cpsVH protein           ORF01310         SAG1169         163         CpsVG           ORF01311         SAG1170         149         CpsF           ORF01312         SAG1171         462         CpsE           ORF01313         SAG1172         229         CpsD protein           ORF01314         SAG1173         230         cpsC protein           ORF01315         SAG1174         243         capsular polysaccharide biosynthesis protein CpsE           ORF01316         SAG1175         485         capsular polysaccharide synthesis operon transcriptional regulator CpsY           ORF01318         SAG1177         255         cpsIaS protein           ORF01319         SAG1178         236         purine nucleoside phosphorylase		SAG1164	321	cpsVJ protein
ORF01308 SAG1167 241 cpsVM protein ORF01309 SAG1168 364 cpsVH protein ORF01310 SAG1169 163 CpsVG ORF01311 SAG1170 149 CpsF ORF01312 SAG1171 462 CpsE ORF01313 SAG1172 229 CpsD protein ORF01314 SAG1173 230 cpsC protein ORF01315 SAG1174 243 capsular polysaccharide biosynthesis protein CpsE ORF01316 SAG1175 485 capsular polysaccharide biosynthesis protein CpsE ORF01317 SAG1176 290 capsular polysaccharide synthesis operon transcriptional regulator CpsY ORF01318 SAG1177 255 cpsIaS protein ORF01319 SAG1178 236 purine nucleoside phosphorylase		SAG1165	327	cpsVO protein
ORF01309 SAG1168 364 cpsVH protein  ORF01310 SAG1169 163 CpsVG  ORF01311 SAG1170 149 CpsF  ORF01312 SAG1171 462 CpsE  ORF01313 SAG1172 229 CpsD protein  ORF01314 SAG1173 230 cpsC protein  ORF01315 SAG1174 243 capsular polysaccharide biosynthesis protein CpsE  ORF01316 SAG1175 485 capsular polysaccharide biosynthesis protein CpsE  ORF01317 SAG1176 290 capsular polysaccharide synthesis operon transcriptional regulator CpsY  ORF01318 SAG1177 255 cpsIaS protein  ORF01319 SAG1178 236 purine nucleoside phosphorylase	ORF01307	SAG1166	295	cpsVN protein
ORF01310 SAG1169 163 CpsVG ORF01311 SAG1170 149 CpsF ORF01312 SAG1171 462 CpsE ORF01313 SAG1172 229 CpsD protein ORF01314 SAG1173 230 cpsC protein ORF01315 SAG1174 243 capsular polysaccharide biosynthesis protein CpsE ORF01316 SAG1175 485 capsular polysaccharide biosynthesis protein CpsE ORF01317 SAG1176 290 capsular polysaccharide synthesis operon transcriptional regulator CpsY ORF01318 SAG1177 255 cpsIaS protein ORF01319 SAG1178 236 purine nucleoside phosphorylase	ORF01308	SAG1167	241	cpsVM protein
ORF01311 SAG1170 149 CpsF ORF01312 SAG1171 462 CpsE ORF01313 SAG1172 229 CpsD protein ORF01314 SAG1173 230 cpsC protein ORF01315 SAG1174 243 capsular polysaccharide biosynthesis protein CpsE ORF01316 SAG1175 485 capsular polysaccharide biosynthesis protein CpsE ORF01317 SAG1176 290 capsular polysaccharide synthesis operon transcriptional regulator CpsY ORF01318 SAG1177 255 cpsIaS protein ORF01319 SAG1178 236 purine nucleoside phosphorylase	ORF01309	SAG1168	364	cpsVH protein
ORF01312 SAG1171 462 CpsE ORF01313 SAG1172 229 CpsD protein ORF01314 SAG1173 230 cpsC protein ORF01315 SAG1174 243 capsular polysaccharide biosynthesis protein CpsE ORF01316 SAG1175 485 capsular polysaccharide biosynthesis protein CpsE ORF01317 SAG1176 290 capsular polysaccharide synthesis operon transcriptional regulator CpsY ORF01318 SAG1177 255 cpsIaS protein ORF01319 SAG1178 236 purine nucleoside phosphorylase	ORF01310	SAG1169	163	CpsVG
ORF01312 SAG1171 462 CpsE ORF01313 SAG1172 229 CpsD protein ORF01314 SAG1173 230 cpsC protein ORF01315 SAG1174 243 capsular polysaccharide biosynthesis protein CpsE ORF01316 SAG1175 485 capsular polysaccharide biosynthesis protein CpsA ORF01317 SAG1176 290 capsular polysaccharide synthesis operon transcriptional regulator CpsY ORF01318 SAG1177 255 cpsIaS protein ORF01319 SAG1178 236 purine nucleoside phosphorylase	ORF01311	SAG1170	149	CpsF
ORF01313 SAG1172 229 CpsD protein ORF01314 SAG1173 230 cpsC protein ORF01315 SAG1174 243 capsular polysaccharide biosynthesis protein CpsE ORF01316 SAG1175 485 capsular polysaccharide biosynthesis protein CpsE ORF01317 SAG1176 290 capsular polysaccharide synthesis operon transcriptional regulator CpsY ORF01318 SAG1177 255 cpsIaS protein ORF01319 SAG1178 236 purine nucleoside phosphorylase	ORF01312	SAG1171	462	- <del> </del>
ORF01314 SAG1173 230 cpsC protein ORF01315 SAG1174 243 capsular polysaccharide biosynthesis protein CpsE ORF01316 SAG1175 485 capsular polysaccharide biosynthesis protein CpsE ORF01317 SAG1176 290 capsular polysaccharide synthesis operon transcriptional regulator CpsY ORF01318 SAG1177 255 cpsIaS protein ORF01319 SAG1178 236 purine nucleoside phosphorylase	ORF01313			CpsD protein
ORF01315 SAG1174 243 capsular polysaccharide biosynthesis protein CpsE ORF01316 SAG1175 485 capsular polysaccharide biosynthesis protein CpsE ORF01317 SAG1176 290 capsular polysaccharide synthesis operon transcriptional regulator CpsY ORF01318 SAG1177 255 cpsIaS protein ORF01319 SAG1178 236 purine nucleoside phosphorylase	ORF01314			
ORF01316 SAG1175 485 capsular polysaccharide biosynthesis protein CpsA ORF01317 SAG1176 290 capsular polysaccharide synthesis operon transcriptional regulator CpsY ORF01318 SAG1177 255 cpsIaS protein ORF01319 SAG1178 236 purine nucleoside phosphorylase				capsular polysaccharide biosynthesis protein CpsB
ORF01317 SAG1176 290 capsular polysaccharide synthesis operon transcriptional regulator CpsY ORF01318 SAG1177 255 cpslaS protein ORF01319 SAG1178 236 purine nucleoside phosphorylase	00564646			
transcriptional regulator CpsY ORF01318 SAG1177 255 cpslaS protein ORF01319 SAG1178 236 purine nucleoside phosphorylase	OKF01316	SAG1175	485	capsular polysaccharide biosynthesis protein CpsA
transcriptional regulator CpsY ORF01318 SAG1177 255 cpslaS protein ORF01319 SAG1178 236 purine nucleoside phosphorylase	ORF01317	SAG1176	290	capsular polysaccharide synthesis operon
ORF01318 SAG1177 255 cpslaS protein ORF01319 SAG1178 236 purine nucleoside phosphorylase				
ORF01319 SAG1178 236 purine nucleoside phosphorylase		SAG1177	255	
	ORF01319	SAG1178	4	
	ORF01320			voltage-gated chlorid channel family protein, putative
ORF01321 SAG1180 269 purine nucleoside phosphorylas	ORF01321	SAG1180	269	purine nucleoside phosphorylas
ORF01322 SAG1181 135 arsenate reductase	ORF01322	SAG1181	135	
ORF01323 SAG1182 403 phosphopentomutase	ORF01323			
ORF01324 SAG1183 223 ribose 5-phosphate isomerase	ORF01324			



ORF Ref No.	SAGxxxx Ref No.	aa	Annotation
ORF01326	SAG1184	236	conserved hypothetical protein
ORF01327	SAG1185	262	tributyrin esterase
ORF01328	SAG1186	553	metallo-beta-lactamase superfamily protein
ORF01329	SAG1187	253	ABC transporter, ATP-binding protein
ORF01330	SAG1188	287	ABC transporter, permease protein
ORF01331	SAG1189	334	conserved hypothetical protein
ORF01332	SAG1190	551	adherence and virulence protein A
ORF01333	SAG1191	239	alpha-acetolactate decarboxylase
ORF01334	SAG1192	560	acetolactate synthase, catabolic
ORF01335	SAG1193	408	TPR domain protein
ORF01336	SAG1194	396	membrane protein
ORF01337	SAG1195	153	MutT/nudix family protein
ORF01338	SAG1196	160	mutator MutT protein
ORF01339	SAG1197	1072	
ORF01340	SAG1197 SAG1198		hyaluronidase
ORF01341		348	dTDP-glucose 4,6-dehydratase
ORF01341	SAG1199	197	dTDP-4-dehydrorhamnose 3,5-epimerase
	SAG1200	289	glucose-1-phosphate thymidylyltransferase
ORF01343	SAG1201	367	iminodiacetate oxidase, putative
ORF01344	SAG1202	262	conserved hypothetical protein TIGR00486
ORF01345	SAG1203	227	conserved hypothetical protein
ORF01346	SAG1204	226	DNA replication protein Dnad, putative
ORF01347	SAG1205	172	adenine phosphoribosyltransferase
ORF01348	SAG1206	854	conserved domain protein
ORF01349	SAG1207	32	hypothetical protein
ORF01350	SAG1208	732	single-stranded-DNA-specific exonuclease RecJ
ORF01351	SAG1209	253	oxidoreductase, short chain dehydrogenase/reductase family
ORF01352	SAG1210	309	metallo-beta-lactamase superfamily protein
ORF01353	SAG1211	215	conserved hypothetical protein
ORF01354	SAG1212	412	GTP-binding protein HflX
ORF01355	SAG1213	296	tRNA delta(2)-isopentenylpyrophosphate transferase
ORF01356	SAG1214	58	hypothetical protein
ORF01357	SAG1215	305	exfoliative toxin A, putative
ORF01358	SAG1216	1252	pullulanase, putative
ORF01361	SAG1217		conserved hypothetical protein, FRAMESHIFT
ORF01362	SAG1218	194	conserved hypothetical protein
ORF01363	SAG1219	468	peptidase, M20/M25/M40 family
ORF01364	SAG1220	200	nitroreductase family protein
ORF01365	SAG1221	<del> </del>	glycerophosphoryl diester phosphodiesterase,
ORF01367	SAG1222	502	putative, POINT MUTATION
ORF01368	SAG1222 SAG1223	593	excinuclease ABC, C subunit
ORF01369		255	conserved hypothetical protein
ORF01369	SAG1224	446	MATE efflux family protein
	SAG1225	136	conserved hypothetical protein
ORF01371	SAG1226	165	conserved hypothetical protein
ORF01372	SAG1227	198	conserved hypothetical protein
ORF01373	SAG1228	96	ISSdy1, transposase OrfA
ORF01374	SAG1229	259	ISSdy1, transposase OrfB
ORF01375	SAG1230	96	conserved hypothetical protein
ORF01377	SAG1231		transposase OrfB, IS3 family, degenerate FRAMESHIFT
ORF01379	SAG1232	77	transposase OrfB, IS3 family, truncation
ORF01380	SAG1233	822	streptococcal histidine triad family protein
ORF01381	SAG1234	306	laminin-binding surface protein
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### Tabl 32: Conv rsi n f ORF Ref N . with SAG Ref Nos.

ORFR fN.	SAGxxxx Ref No.	aa	Ann tation
ORF01382	SAG1235	425	GBSi1, group II intron, maturase
ORF01383	SAG1236		c5a peptidase precursor FRAMESHIFT
ORF01384	SAG1237	444	hypothetical protein
ORF01385	SAG1238	202	hypothetical protein
ORF01386	SAG1239	76	conserved hypothetical protein
ORF01387	SAG1240	125	conserved hypothetical protein, truncation
ORF01388	SAG1241	78	transposase OrfA, IS3 family
ORF01389	SAG1242	67	transposase OrfB, IS3 family, truncation
ORF01390	SAG1243	96	ISSdy1, transposase OrfA FRAMESHIFT
ORF01391	SAG1244	259	ISSdy1, transposase OrfB
ORF01392	SAG1245	38	hypothetical protein
ORF01393	SAG1246	389	hypothetical protein
ORF01394	SAG1247	399	integrase, phage family
ORF01395	SAG1248	75	conserved hypothetical protein
ORF01396	SAG1249	74	transcriptional regulator, Cro/CI family
ORF01397	SAG1250	621	Tn5252, relaxase
ORF01398	SAG1251	121	Tn5252, Orf 9 protein
ORF01399	SAG1252	120	Tn5252, Orf 10 protein
ORF01401	SAG1253	435	transposase, ISL3 family
ORF01403	SAG1254	546	mercuric reductase
ORF01404	SAG1255	130	
31 31-131	UNG1255	130	mercuric resistance operon regulatory protein MerR
ORF01406	SAG1256	142	IS861, transposase OrfB, truncation
ORF01407	SAG1257	709	cation-transporting ATPase, E1-E2 family
ORF01408	SAG1258	122	cadmium efflux system accessory protein
ORF01409	SAG1259	99	conserved hypothetical protein
ORF01410	SAG1260	262	hypothetical protein
ORF01411	SAG1261	198	conserved hypothetical protein
ORF01412	SAG1262	695	cation-transporting ATPase, E1-E2 family
ORF01414	SAG1263		conserved domain protein, FRAMESHIFT
ORF01415	SAG1264	148	transcriptional repressor CopY, putative
ORF01416	SAG1265	206	cadmium resistance transporter, putative
ORF01417	SAG1266	152	hypothetical protein
ORF01418	SAG1267	108	hypothetical protein
ORF01419	SAG1268	230	repressor protein, putative
ORF01420	SAG1269	44	hypothetical protein
ORF01421	SAG1270	471	ImpB/MucB/SamB family protein
ORF01423	SAG1271	116	conserved hypothetical protein
ORF01424	SAG1272	102	conserved hypothetical protein
ORF01425	SAG1273	118	conserved hypothetical protein
ORF01426	SAG1274	129	conserved hypothetical protein
ORF01427	SAG1275	75	hypothetical protein
ORF01428	SAG1276	358	conserved hypothetical protein
ORF01430	SAG1277	163	hypothetical protein
ORF01431	SAG1278	96	hypothetical protein
ORF01432	SAG1279	99	conserved domain protein
ORF01433	SAG1280	2274	Helicases conserved C-terminal domain protein
ORF01434	SAG1281	183	hypothetical protein
ORF01435 ORF01436	SAG1282	63	lipoprotein, putative
ORF01436	SAG1283	1631	cell wall surface anchor family protein
ORF01437	SAG1284	196	abortiv infection protein AbiGI
ORF01439	SAG1285 SAG1286	281	abortive infection protein AbiGII
ORF01440	SAG1286 SAG1287	933	conserved hypothetical protein
ORF01441	SAG1287	776	conserved hypothetical protein
	UAG 1200	117	conserved hypothetical protein, DEGENERATE

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Table 32: C nversion of ORF R f N	s. with SAG R f Nos.

ORF Ref No.	SAGxxxx Ref No.	aa	Annotati n
ORF01442	SAG1289	284	conserved hypothetical protein
ORF01443	SAG1290	80	hypothetical protein
ORF01444	SAG1291	605	Tn5252, Orf 21 protein, internal deletion
ORF01445	SAG1292	162	hypothetical protein
ORF01446	SAG1293	194	protease, putative
ORF01447	SAG1294	77	conserved hypothetical protein
ORF01449	SAG1295	127	conserved hypothetical protein
ORF01450	SAG1295		
ORF01451	SAG1297	142 451	conserved hypothetical protein
3/1/01401	UNG 1231	401	type II DNA modification methyltransferase Spn5252IP
ORF01452	SAG1298	31	hypothetical protein
ORF01453	SAG1299	272	conserved hypothetical protein
ORF01454	SAG1300	57	conserved hypothetical protein
ORF01455	SAG1301	121	ribosomal protein L7/L12
ORF01456	SAG1302	166	ribosomal protein L10
ORF01458	SAG1303	702	ATP-dependent Clp protease, ATP-binding subunit
		, 02	ATP-billing subutilit
ORF01459	SAG1304	32	hypothetical protein
ORF01460	SAG1305	314	homocysteine S-methyltransferase MmuM, putative
ORF01461	SAG1306	458	amino acid permease
ORF01463	SAG1307	216	hypothetical protein
ORF01464	SAG1308	167	hypothetical protein
ORF01465	SAG1309	30	hypothetical protein
ORF01466	SAG1310	182	transcriptional regulator, TetR family
ORF01467	SAG1311	198	GTP-binding protein
ORF01468	SAG1312	408	ATP-dependent Clp protease, ATP-binding subunit
			ClpX
ORF01469	SAG1313	56	conserved hypothetical protein
ORF01470	SAG1314	164	dihydrofolate reductase
ORF01471	SAG1315	279	thymidylate synthase
ORF01472	SAG1316	390	HMG-CoA synthase
ORF01473	SAG1317	427	3-hydroxy-3-methylglutaryl-CoA reductase
ORF01474	SAG1318	149	conserved hypothetical protein
ORF01475	SAG1319	187	hemolysin III, putative
ORF01476	SAG1320	304	conserved hypothetical protein TIGR00147
ORF01477	SAG1321	284	glutathione S-transferase family protein
ORF01478	SAG1322	72	conserved domain protein
ORF01479	SAG1323	331	isopentenyl-diphosphate delta-isomerase
ORF01480	SAG1324	330	phosphomevalonate kinase
ORF01481	SAG1325	314	diphosphomevalonate decarboxylase
ORF01482	SAG1326	292	mevalonate kinase, putative
ORF01483	SAG1327	409	sensor histidine kinase
ORF01484	SAG1328	228	DNA-binding response regulator
ORF01485	SAG1329	208	GTP pyrophosphokinase family protein
ORF01486	SAG1330	68	hypothetical protein
ORF01488	SAG1331	979	R5 protein
ORF01489	SAG1332	146	transcriptional regulator, MarR family, putative
ORF01490	SAG1333	690	5'-nucleotidase family protein
ORF01491	SAG1334	136	polypeptide deformylase, putative
ORF01492	SAG1335	449	NADP-specific glutamate dehydrogenase
ORF01494	SAG1336	169	conserved hypothetical protein
ORF01495	SAG1337		
ORF01496	SAG1337 SAG1338	589	ABC transporter, ATP-binding/permease protein
ORF01497	SAG1338	579	ABC transporter, ATP-binding/permease protein
	OUG 1998	157	acetyltransferase, GNAT family

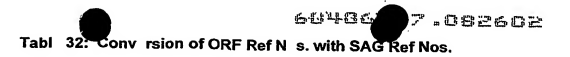


### Table 32: C nversion of ORF R f Nos. with SAG Ref Nos.

ORF R f No.	SAGxxxx Ref No.	aa	Annotation
ORF01498	SAG1340	622	ABC transporter, ATP-binding protein
ORF01499	SAG1341	402	polyA polymerase family protein
ORF01500	SAG1342	282	DegV family protein
ORF01501	SAG1343	126	conserved hypothetical protein
ORF01502	SAG1344	177	hypothetical protein
ORF01503	SAG1345	164	conserved hypothetical protein
ORF01504	SAG1346	641	PTS system, fructose specific IIABC components
ORF01505	SAG1347	303	1-phosphofructokinase
ORF01506	SAG1348	247	lactose phosphotransferase system repressor
ORF01507	SAG1349	411	beta-lactam resistance factor
ORF01508	SAG1350	544	surface antigen-related protein
ORF01509	SAG1351	307	2-dehydropantoate 2-reductase, putative
ORF01510	SAG1352	356	regulatory protein, putative
ORF01511	SAG1353	330	pyridine nucleotide-disulphide oxidoreductase family protein
ORF01512	SAG1354	251	tRNA (guanine-N1)-methyltransferase
ORF01513	SAG1355	172	16S rRNA processing protein RimM
ORF01515	SAG1356	503	transcriptional regulator, RofA family
ORF01516	SAG1357	80	KH domain protein
ORF01517	SAG1358	90	ribosomal protein S16
ORF01518	SAG1359	415	permease, putative
ORF01519	SAG1360	236	ABC transporter, ATP-binding protein
ORF01520	SAG1361	414	conserved hypothetical protein
ORF01522	SAG1362	532	carbamoyl-phosphate synthase, large subunit, putative
ORF01523	SAG1363	356	carbamoyl-phosphate synthase, small subunit
ORF01524	SAG1364	173	pyrimidine operon regulatory protein
ORF01525	SAG1365	296	ribosomal large subunit pseudouridine synthase, RluD subfamily
ORF01526	SAG1366	154	lipoprotein signal peptidase
ORF01527	SAG1367	301	transcriptional regulator, LysR family
ORF01528	SAG1368	94	ribosomal protein L27
ORF01529	SAG1369	112	conserved hypothetical protein .
ORF01530	SAG1370	104	ribosomal protein L21
ORF01531	SAG1371	392	conserved hypothetical protein
ORF01532	SAG1372	404	thiamine biosynthesis protein Thil
ORF01533	SAG1373	381	cysteine desulphurase
ORF01535	SAG1374	150	conserved hypothetical protein
ORF01536	SAG1375	449	glutathione reductase
ORF01537	SAG1376	111	conserved hypothetical protein
ORF01538	SAG1377	388	chorismate synthase
ORF01539	SAG1378	355	3-dehydroquinate synthase
ORF01540	SAG1379	225	3-dehydroquinate dehydratase
ORF01541	SAG1380	385	conserved hypothetical protein
ORF01542	SAG1381	714	sulfatase
ORF01543	SAG1382	119	ribosomal protein L20
ORF01544	SAG1383	66	ribosomal protein L35
ORF01545	SAG1384	176	translation initiation factor IF-3
ORF01546	SAG1385	227	cytidylate kinase
ORF01547	SAG1386	174	conserved hypothetical protein
ORF01548	SAG1387	65	ferredoxin, 4Fe-4S
ORF01549	SAG1388	163	conserved hypothetical protein
ORF01550	SAG1389	406	peptidase t
ORF01551	SAG1390	544	polysaccharide biosynthesis protein, putative



alanyl-D-glutamate2,6-
nsporter, ATP-binding protein
nsporter, substrate-binding
nsporter, permease protein
nsporter, permease protein
protein
tase, manganese-dependent
-activating enzyme
protein
protein TIGR01212
tative
r family protein
r family protein
r family protein
or, RofA family FRAMESHIFT
roup 1 family protein
synthesis protein, putative
synthesis protein, putative
roup 2 family protein
roup 2 family protein
nydratase, putative
-methyl-D-erythritol synthase,
al protein
group 2 family protein
group 2 family protein
al protein
nose reductase
al protein
ma-70 factor
echanosensitive channel protein
al protein ,
sporter, amino acid-binding
er family protein
al protein
tein



ORF Ref N .	SAGxxxx Ref No.	aa	Annotation
ORF01601	SAG1437	55	hypothetical protein
ORF01602	SAG1438	754	glycogen phosphorylase
ORF01603	SAG1439	498	4-alpha-glucanotransferase
ORF01604	SAG1440	342	maltose operon repressor MalR, putative
ORF01605	SAG1441	415	maltose/maltodextrin ABC transporter,
		'''	maltose/maltodextrin-binding protein
ORF01606	SAG1442	456	maltose ABC transporter, permease protein
ORF01607	SAG1443	278	maltose ABC transporter, permease protein
ORF01608	SAG1444	490	proton/peptide symporter family protein
ORF01610	SAG1445		MutT/nudix family protein, FRAMESHIFT
ORF01611	SAG1446	62	hypothetical protein
ORF01612	SAG1447	441	conserved hypothetical protein
ORF01613	SAG1448	502	glycosyl transferase, group 1 family protein
ORF01614	SAG1449	795	preprotein translocase SecA subunit, putative
ORF01615	SAG1450	330	conserved domain protein
ORF01617	SAG1451	494	conserved hypothetical protein
ORF01618	SAG1452	514	conserved hypothetical protein
ORF01619	SAG1453	409	preprotein translocase SecY family protein
ORF01621	SAG1454	398	conserved hypothetical protein
ORF01622	SAG1455	295	glycosyl transferase, group 2 family protein
ORF01623	SAG1456	312	glycosyl transferase, family 8, degenerate
ORF01624	SAG1457	129	IS1381, transposase OrfB
ORF01625	SAG1458	127	IS1381, transposase OrfA
ORF01626	SAG1459	413	glycosyl transferase family 8
ORF01627	SAG1460	401	glycosyl transferase, family 8
ORF01628	SAG1461	335	conserved hypothetical protein
ORF01630	SAG1462	970	cell wall surface anchor family protein
ORF01632	SAG1463		transcriptional regulator, RofA family POINT MUTATION
ORF01634	SAG1464	663	excinuclease ABC, B subunit
ORF01635	SAG1465	306	protease, putative
ORF01636	SAG1466	727	glutamine ABC transporter, glutamine-binding
	-		protein/permease protein, putative
ORF01637	SAG1467	246	glutamine ABC transporter, ATP-binding protein, GlnQ putative
ORF01638	SAG1468	116	conserved hypothetical protein
ORF01639	SAG1469	52	conserved hypothetical protein
ORF01640	SAG1470	437	GTP-binding protein, GTP1/Obg family
ORF01641	SAG1471	42	conserved hypothetical protein
ORF01643	SAG1472	413	aminopeptidase PepS
ORF01645	SAG1473	192	cell wall surface anchor family protein
ORF01646	SAG1474	680	amidase family protein
ORF01647	SAG1475	240	ribosomal small subunit pseudouridine synthase A
ORF01648	SAG1476	280	oxidoreductase, aldo/keto reductase family
ORF01650	SAG1477	224	nitroreductase family protein
ORF01651	SAG1478	130	lactoylglutathione lyase
ORF01652	SAG1479	308	glycosyl transferase, group 2 family protein
ORF01653	SAG1480	462	amino acid permease
ORF01654	SAG1481	155	SsrA-binding protein
ORF01655	SAG1482	801	exoribonuclease, VacB/Rnb family
ORF01657	SAG1483	78	preprotein translocase, SecG subunit
ORF01658	SAG1485	389	multi-drug resistance protein
ORF01658 ORF01660	SAG1485 SAG1486	389 548	multi-drug resistance protein hypothetical protein

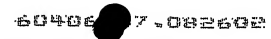
### Table 32: Conversion of ORFR fN s. with SAGR fNos.

ORF Ref N .	SAGxxxx Ref No.	aa	Annotati n
ORF01662	SAG1488	195	dephospho-CoA kinase
ORF01663	SAG1489	273	formamidopyrimidine-DNA glycosylase
ORF01665	SAG1490	282	transcriptional regulator, MutR family
ORF01666	SAG1491	530	hypothetical protein
ORF01667	SAG1492	58	hypothetical protein
ORF01668	SAG1493		hypothetical protein
ORF01670	SAG1494	32	hypothetical protein
ORF01672	SAG1495	81	protease, putative, POINT MUTATION
ORF01673	SAG1496	110	hypothetical protein
ORF01674	SAG1497	37	hypothetical protein
ORF01675	SAG1498	133	hypothetical protein
ORF01677	SAG1499	299	GTP-binding protein Era
ORF01678	SAG1500	132	diacylglycerol kinase
ORF01679	SAG1501	161	conserved hypothetical protein TIGR00043
· ORF01680	SAG1502	268	tetracenomycin polyketide synthesis O-
			methyltransferase TcmP, putative
ORF01681	SAG1503	39	hypothetical protein
ORF01682	SAG1504	38	hypothetical protein
ORF01683	SAG1505	158	MutT/nudix family protein
ORF01684	SAG1506	267	hypothetical protein
ORF01685	SAG1507	345	PhoH family protein
ORF01686	SAG1508	590	67 kDa Myosin-crossreactive streptococcal antigen
			The my com-crossreactive streptococcar antiger
ORF01687	SAG1509	71	conserved hypothetical protein
ORF01688	SAG1510	169	peptide methionine sulfoxide reductase
ORF01689	SAG1511	284	conserved hypothetical protein
ORF01690	SAG1512	185	ribosome recycling factor
ORF01691	SAG1513	242	uridylate kinase
ORF01692	SAG1514	226	peptide ABC transporter, ATP-binding protein
ORF01693	SAG1515	262	peptide ABC transporter, ATP-binding protein
ORF01694	SAG1516	255	peptide ABC transporter, permease protein
ORF01695	SAG1517	314	peptide ABC transporter, permease protein
ORF01696	SAG1518	525	peptide ABC transporter, peptide-binding protein
ORF01697	SAG1519	229	ribosomal protein L1
ORF01698	SAG1520	141	ribosomal protein L11
ORF01699	SAG1521	388	transposase, IS30 family, putative
ORF01700	SAG1522	460	transporter, major facilitator family
ORF01702	SAG1523	404	peptidase, M20/M25/M40 family
ORF01703	SAG1524	294	transcriptional regulator, LysR family
ORF01704	SAG1525	117	conserved hypothetical protein
ORF01705	SAG1526	178	IS861, transposase OrfA
ORF01706	SAG1527	277	IS861, transposase OrfB
ORF01707	SAG1528	571	chorismate binding enzyme
ORF01708	SAG1529	. 785	FtsK/SpollIE family protein
ORF01709	SAG1530	267	peptidyl-prolyl cis-trans isomerase, cyclophilin-type
			The state of the s
ORF01710	SAG1531	277	manganese ABC transporter, permease protein
ORF01711	SAG1532	238	manganese ABC transporter, ATP-binding protein
ORF01712	SAG1533	308	manganese ABC transporter, manganese-binding
			adhesion liprotein
ORF01713	SAG1534	215	iron-dependent transcriptional regulator
ORF01714	SAG1535	229	5-methylthioadenosine nucleosidase/S-
0550			adenosylhomocysteine nucleosidase
ORF01715	SAG1536	89	conserved hypothetical protein
ORF01716	SAG1537	184	MutT/nudix family protein

# Table 32: Conv rsi n of ORF R f Nos. with SAG Ref Nos.

ORF Ref No.	SAGxxxx R fN .	aa	Annotati n
ORF01718	SAG1538	459	UDP-N-acetylglucosamine pyrophosphorylase
ORF01719	SAG1539	31	hypothetical protein
ORF01720	SAG1540	137	conserved hypothetical protein
ORF01721	SAG1541	125	glyoxalase family protein
ORF01722	SAG1542	318	oxidoreductase, Gfo/Idh/MocA family
ORF01724	SAG1543	0.0	conserved hypothetical protein, FRAMESHIFT
ORF01725	SAG1544	232	gluconate 5-dehydrogenase, putative
ORF01726	SAG1545	78	conserved hypothetical protein
ORF01727	SAG1546	82	conserved hypothetical protein
ORF01729	SAG1547	166	acetyltransferase, GNAT family
ORF01730	SAG1548	422	glycosyl transferase, group 2 family protein
ORF01731	SAG1549	127	
ORF01731	SAG1549	129	IS1381, transposase OrfA IS1381, transposase OrfB
ORF01733	SAG1550	67	
ORF01733	SAG1551		hypothetical protein
ORF01735	SAG1552	719 477	conserved hypothetical protein
ORF01736	SAG1553	225	hypothetical protein
			hypothetical protein
ORF01737 ORF01738	SAG1555	231	hypothetical protein
ORF01738	SAG1556	445	branched-chain amino acid transport system II carrier protein
ORF01739	SAG1557	665	methionyl-tRNA synthetase
ORF01740	SAG1558	291	tellurite resistance protein TehB
ORF01741	SAG1559	231	membrane protein, putative
ORF01742	SAG1560	40	hypothetical protein
ORF01743	SAG1560	405	PTS system component, putative
ORF01744	SAG1562	280	conserved hypothetical protein
ORF01745	SAG1563	275	exodeoxyribonuclease
ORF01746	SAG1564	118	conserved hypothetical protein
ORF01747	SAG1565	158	methylated-DNA-protein-cysteine S-methyltransferase
0	0,101000	100	inetrylated-DNA-protein-cysteine 5-metryluansierase
ORF01748	SAG1566	393	D-isomer specific 2-hydroxyacid dehydrogenase family
		1	protein
ORF01749	SAG1567	182	acetyltransferase, GNAT family
ORF01750	SAG1568		phosphoserine aminotransferase FRAMESHIFT
ORF01752	SAG1569	211	copper homeostasis protein CutC, putative
ORF01753	SAG1570	34	conserved hypothetical protein
ORF01754	SAG1571	53	hypothetical protein
ORF01755	SAG1572	287	tetrapyrrole methylase family protein
ORF01756	SAG1573	108	conserved hypothetical protein
ORF01758	SAG1574	287	DNA polymerase III, delta prime subunit, putative
ORF01759	SAG1575	211	thymidylate kinase
ORF01761	SAG1576	267	transposase, IS30 family, putative, truncation
ORF01763	SAG1577	219	AcuB family protein
ORF01764	SAG1578	236	branched-chain amino acid ABC transporter, ATP-
}			binding protein
ORF01765	SAG1579	254	branched-chain amino acid ABC transporter, ATP-
			binding protein
ORF01766	SAG1580	317	branched-chain amino acid ABC transporter,
			permease protein
ORF01767	SAG1581	289	branched-chain amino acid ABC transporter,
			permease protein
ORF01769	\$AG1582	388	branched-chain amino acid ABC transporter, amino
00504555	-		acid-binding protein
ORF01770	SAG1583	81	conserved hypothetical protein
ORF01772	SAG1584	377	IS1548, transposase





# Table 32: C nv rsion of ORF R f Nos. with SAG R f Nos.

ORF Ref No.	SAGxxxx Ref N .	aa	Annotation
ORF01773	SAG1585	196	ATP-dependent Clp protease, proteolytic subunit ClpP
ORF01774	SAG1586	209	uracil phosphoribosyltransferase
ORF01775	SAG1587	389	aminotransferase, class I
ORF01777	SAG1588	182	RNA methyltransferase, TrmH family, group 2
ORF01778	SAG1589	450	amino acid permease, putative
ORF01779	SAG1590	449	potassium uptake protein, Trk family
ORF01780	SAG1591	475	cation uptake protein, Trk family
ORF01781	SAG1592	83	conserved hypothetical protein TIGR00278
ORF01782	SAG1593	240	ribosomal large subunit pseudouridine synthase B
ORF01783	SAG1594	194	conserved hypothetical protein TIGR00281
ORF01784	SAG1595	235	Uncharacterized ACR, COG1354
ORF01785	SAG1596	246	integrase/recombinase, phage integrase family
ORF01786	SAG1597	157	CBS domain protein
ORF01787	SAG1598	173	conserved hypothetical protein
ORF01788	SAG1599	324	HAM1 protein
ORF01789	SAG1600	264	glutamate racemase
ORF01790	SAG1601	79	conserved hypothetical protein
ORF01791	SAG1602	180	membrane protein, putative
ORF01792	SAG1603	173	transcriptional regulator, biotin repressor family
ORF01793	SAG1604	229	membrane protein, putative
ORF01794	SAG1605	167	conserved hypothetical protein
ORF01795	SAG1606	247	RNA methyltransferase, TrmH family
ORF01796	SAG1607	92	acylphosphatase
ORF01797	SAG1608	310	membrane protein, putative
ORF01799	SAG1609	221	amino acid ABC transporter, permease protein
ORF01800	SAG1610	285	amino acid ABC transporter, substrate-binding protein
ORF01801	SAG1611	486	amidase family protein
ORF01802	SAG1612	160	transcription elongation factor GreA
ORF01803	SAG1613	600	Uncharacterized BCR, YceG family COG1559, putative
ORF01804	SAG1614	167	acetyltransferase, GNAT family
ORF01805	SAG1615	443	UDP-N-acetylmuramatealanine ligase
ORF01806	SAG1616	205	conserved hypothetical protein
ORF01807	SAG1617	32	hypothetical protein
ORF01808	SAG1618	1032	Snf2 family protein
ORF01810	SAG1619	377	IS1548, transposase
ORF01811	SAG1620	436	phosphoglycerate dehydrogenase-related protein
ORF01812	SAG1621	300	primosomal protein Dnal
ORF01813	SAG1622	391	conserved hypothetical protein
ORF01814	SAG1623	159	conserved hypothetical protein TIGR00244
ORF01815	SAG1624	501	sensor histidine kinase CsrS
ORF01816	SAG1625	229	DNA-binding response regulator CsrR
ORF01817	SAG1626	177	conserved hypothetical protein
ORF01818	SAG1627	296	heat shock protein HtpX
ORF01820	SAG1628	184	lemA protein
ORF01821	SAG1629	237	glucose-inhibited division protein B
ORF01822	SAG1630	459	sodium transport family protein
ORF01823	SAG1631	223	potassium uptake protein, Trk family, putativ
ORF01824	SAG1632	276	cobalt transport family protein
ORF01825	SAG1633	558	ABC transporter, ATP-binding protein
ORF01826	SAG1634	212	conserved hypothetical protein





## Table 32: Conv rsion fORF Ref Nos. with SAG R f Nos.

ORF Ref N .	SAGxxxx Ref No.	aa	Annotati n
ORF01828	SAG1636	455	branched-chain amino acid transport system II carrier
			protein
ORF01829	SAG1637	351	alcohol dehydrogenase, zinc-containing
ORF01830	SAG1638	230	ABC transporter, permease protein
ORF01831	SAG1639	356	ABC transporter, ATP-binding protein
ORF01832	SAG1640	458	peptidase, M20/M25/M40 family
ORF01833	SAG1641	274	lipoprotein, putative
ORF01834	SAG1642	277	ABC transporter, substrate-binding protein
ORF01835	SAG1643	229	glutamine amidotransferase, class i
ORF01836	SAG1644	37	hypothetical protein
ORF01837	SAG1645	238	conserved hypothetical protein TIGR01033
ORF01838	.SAG1646	32	hypothetical protein
ORF01839	SAG1647	328	dihydroxyacetone kinase family protein
ORF01840	SAG1648	178	transcriptional regulator, TetR family, putative
ORF01842	SAG1649	. 37	hypothetical protein
ORF01843	SAG1650	329	dihydroxyacetone kinase family protein
ORF01844	SAG1651	192	dihydroxyacetone kinase family protein
ORF01845	SAG1652	124	conserved hypothetical protein
ORF01846	SAG1653	237	glycerol uptake facilitator protein
ORF01847	SAG1654	134	conserved hypothetical protein
ORF01848	SAG1655	237	transcriptional regulator, MerR family
ORF01849	SAG1656	369	conserved hypothetical protein
ORF01850	SAG1657	83	hypothetical protein
ORF01851	SAG1658	244	conserved hypothetical protein
ORF01852	SAG1659	118	iojap-related protein
ORF01853	SAG1660	173	isochorismatase family protein
ORF01854	SAG1661	195	conserved hypothetical protein TIGR00488
ORF01855	SAG1662	210	conserved hypothetical protein TIGR00482
ORF01856	SAG1663	105	conserved hypothetical protein TIGR00482  conserved hypothetical protein TIGR00253
ORF01857	SAG1664	372	GTP-binding protein
ORF01858	SAG1665	177	
ORF01859	SAG1666	295	hydrolase, haloacid dehalogenase-like family membrane protein
ORF01860	SAG1667	480	
ORF01861	SAG1668	488	glutamyl-tRNA(Gln) amidotransferase, B subunit
ORF01862	SAG1669	100	glutamyl-tRNA(GIn) amidotransferase, A subunit
ORF01863	SAG1670	881	glutamyl-tRNA(GIn) amidotransferase, C subunit
ORF01864	SAG1671	276	pyruvate phosphate dikinase
ORF01865	SAG1672		conserved hypothetical protein
ORF01866	SAG1673	170	CBS domain protein
ORF01867		377	3-hydroxyacyl-CoA dehydrogenase family protein
ORF01869	SAG1674	182	isochorismatase family protein
ORF01870	SAG1675	261	transcriptional regulator CodY, putative
ORF01871	SAG1676	403	arninotransferase, class I
	SAG1677	137	universal stress protein family FRAMESHIFT
ORF01872	SAG1678	460	hydrolase, haloacid dehalogenase-like family
ORF01873	SAG1679	320	asparaginase family protein
ORF01874	SAG1680	292	shikimate 5-dehydrogenase
ORF01875	SAG1681	304	oxidoreductase, aldo/keto reductase family
ORF01876	SAG1682	671	ATP-dependent DNA helicase RecG
ORF01877	SAG1683	512	immunogenic secreted protein, putative
ORF01878	SAG1684	366	alanine racemase
ORF01879	SAG1685	119	holo-(acyl-carrier-protein) synthase
ORF01880	SAG1686	335	phospho-2-dehydro-3-deoxyheptonate aldolase
ORF01881	SAG1687	842	preprotein translocas , SecA subunit
ORF01882	SAG1688	315	mannose-6-phosphat isomerase, class I
ORF01883	SAG1689	293	fructokinase



# Table 32: C nversi n of ORF Ref Nos. with SAG R f Nos.

ORF Ref No.	SAGxxxx Ref No.	aa	Annotation
ORF01885	SAG1690	639	PTS system IIABC components
ORF01886	SAG1691	479	sucrose-6-phosphate hydrolase
ORF01887	SAG1692	320	sucrose operon repressor ScrR
ORF01888	SAG1693	144	N utilization substance protein B
ORF01889	SAG1694	129	conserved hypothetical protein
ORF01890	SAG1695	186	translation elongation factor P
ORF01892	SAG1696	38	hypothetical protein
ORF01893	SAG1697	48	hypothetical protein
ORF01894	SAG1698	99	conserved hypothetical protein
ORF01895	SAG1699	30	hypothetical protein
ORF01896	SAG1700	76	hypothetical protein
ORF01897	SAG1701	56	hypothetical protein
ORF01898	SAG1702	41	hypothetical protein
ORF01899	SAG1703	54	hypothetical protein
ORF01900	SAG1704	150	
ORF01902	SAG1705	130	cytidine/deoxycytidylate deaminase family protein
ORF01903	SAG1706	238	peptidase, M24 family POINT MUTATION
ORF01904	SAG1707	499	conserved hypothetical protein
ORF01905	SAG1708	38	drug resistance transporter, EmrB/QacA family
ORF01906	SAG1709	942	hypothetical protein
ORF01907	SAG1710	223	excinuclease ABC, A subunit
ORF01908	SAG1710		conserved hypothetical protein
ORF01909		314	magnesium transporter, CorA family
ORF01910	SAG1712	79	ribosomal protein S18
ORF01911	SAG1713	163	single-strand binding protein
ORF01912	SAG1714	95	ribosomal protein S6
ORF01912	SAG1715	374	A/G-specific adenine glycosylase
ORF01913	SAG1716	197	transcriptional regulator, Cro/Cl family
ORF01915	SAG1717 SAG1718	104 166	thioredoxin
ORF01916	SAG1718	779	PAP2 family protein
ORF01917	SAG1719	180	MutS2 family protein
ORF01918	SAG1721	103	conserved hypothetical protein
ORF01919	SAG1722	297	conserved hypothetical protein
ORF01920	SAG1723	197	
ORF01921	SAG1724		signal peptidase I
ORF01922	SAG1725	806	helicase, putative
ORF01923	SAG1726	160 364	conserved hypothetical protein
ORF01924	SAG1726		DNA-damage inducible protein P
ORF01925	SAG1727 SAG1728	770	formate acetyltransferase
ORF01926	SAG1728	124	FMN-binding protein
ORF01927		309	conserved hypothetical protein
ORF01927	SAG1730	251	proteinase, putative, degenerate, FRAMESHIFT
ORF01929	SAG1731	298	membrane protein, putative
ORF01929 ORF01930	SAG1732	282	glycerol uptake facilitator protein, putative
ORF01930	SAG1733	150	universal stress protein family
ORF01931	SAG1734	400	transporter, putative
	SAG1735	219	transcriptional regulator, Crp/Fnr family
ORF01933 ORF01934	SAG1736	761	X-pro dipeptidyl-peptidase
	SAG1737	119	hypothetical protein
ORF01936	SAG1738	326	polyprenyl synthetase family protein
ORF01937 ORF01938	SAG1739	582	ABC transporter, ATP-binding protein CydC
	SAG1740	572	ABC transporter, ATP-binding protein CydD
ORF01939	SAG1741	339	cytochrome d ubiquinol oxidase, subunit II
ORF01940	SAG1742	475	cytochrome d oxidase, subunit I
ORF01941	SAG1743	402	pyridine nucleotide-disulphide oxidoreductase family
L	<u> </u>		protein



# Tabl 32: Conversi n of ORF R f Nos. with SAG Ref Nos.

ORF Ref N .	SAGxxxx Ref No.	aa	Annotation
ORF01942	SAG1744	299	prenyltransferas , UbiA family
ORF01943	SAG1745	148	hypothetical protein
ORF01944	SAG1746	35	hypothetical protein
ORF01945	SAG1747	99	conserved hypothetical protein TIGR00103
ORF01946	SAG1748	396	cyclopropane-fatty-acyl-phospholipid synthase
ORF01947	SAG1749	241	transcriptional regulator, merR family
ORF01948	SAG1750	195	exonuclease
ORF01949	SAG1751	178	
ORF01950	SAG1752	375	conserved hypothetical protein conserved hypothetical protein TIGR00275
ORF01951	SAG1753	260	conserved hypothetical protein TIGR00275
ORF01952	SAG1754	89	
ORF01953	SAG1755	38	ribosomal protein S14
ORF01954	SAG1756		hypothetical protein
ORF01957		341	conserved hypothetical protein
ORF01958	SAG1757	336	O-sialoglycoprotein endopeptidase family protein
OKF01936	SAG1758	135	ribosomal-protein-alanine acetyltransferase, putative
ORF01960	SAG1759	230	glycoprotease family protein, putative
ORF01961	SAG1760	76	conserved hypothetical protein
ORF01962	SAG1761	559	metallo-beta-lactamase superfamily protein
ORF01963	SAG1762	169	conserved hypothetical protein
ORF01964	SAG1763	448	glutamine synthetase, type I
ORF01965	SAG1764	123	transcriptional regulator GlnR
ORF01967	SAG1765	179	conserved hypothetical protein
ORF01969	SAG1766	398	phosphoglycerate kinase
ORF01970	SAG1767	289	acid phosphatase
ORF01971	SAG1768	336	glyceraldehyde 3-phosphate dehydrogenase
ORF01972	SAG1769	692	translation elongation factor G
ORF01973	SAG1770	156	ribosomal protein S7
ORF01974	SAG1771	137	ribosomal protein S12
ORF01975	SAG1772	270	pur operon repressor
ORF01976	SAG1773	313	HD domain protein
ORF01977	SAG1774 ·	424	conserved hypothetical protein
ORF01978	SAG1775	210	conserved hypothetical protein
ORF01979	SAG1776	220	ribulose-phosphate 3-epimerase
ORF01980	SAG1777	290	conserved hypothetical protein TIGR00157
ORF01981	SAG1778	283	rRNA (guanine-N1-)-methyltransferase, putative
ORF01983	SAG1779	290	dimethyladenosine transferase
ORF01984	SAG1780	163	hypothetical protein
ORF01985	SAG1781	186	primase-related protein
ORF01987	SAG1782	260	deoxyribonuclease, TatD family
ORF01988	SAG1783	90	hypothetical protein
ORF01989	SAG1784	130	hypothetical protein
ORF01990	SAG1785	430	hypothetical protein
ORF01991	SAG1786	130	
ORF01992	SAG1787	420	hypothetical protein dltD protein
ORF01993	SAG1788	79	D-alanyl carrier protein
ORF01994	SAG1789	421	ditB protein
ORF01996	SAG1789	511	
ORF01997	SAG1791	395	D-alanine-activating enzyme sensor histidine kinase
ORF01998	SAG1791	224	
ORF01999	SAG1792 SAG1793		DNA-binding response regulator
ORF02000	SAG1793	44	ribosomal protein L34
ORF02001	SAG1794 SAG1795	451	membrane protein, putativ
ORF02001		388	transposase, IS30 family, putative
ORF02002	SAG1798	575	amino acid ABC transporter, permease protein
UNI 02004	SAG1797	407	amino acid ABC transporter, ATP-binding protein



# Table 32: Conv rsion of ORF Ref Nos. with SAG R f N s.

ORF Ref N .	SAGxxxx Ref N .	aa	Annotation
ORF02005	SAG1798	39	hypothetical protein
ORF02006	SAG1799	792	xylulose-5-phosphate/fructose-6-phosphate
		1	phosphoketolase
ORF02007	SAG1800	363	conserved hypothetical protein
ORF02008	SAG1801	559	transcriptional antiterminator, BglG family
ORF02009	SAG1802	253	conserved hypothetical protein
ORF02010	SAG1803	505	carbohydrate kinase, FGGY family
ORF02011	SAG1804	329	hypothetical protein
ORF02012	SAG1805	483	PTS system component, putative
ORF02015	SAG1806	318	glyoxylate reductase, NADH-dependent
ORF02016	SAG1807	339	hypothetical protein
ORF02017	SAG1808	327	sugar binding transcriptional regulator, Lacl family
ODE00040	0101000		
ORF02018	SAG1809	215	transaldolase family protein
ORF02019	SAG1810	238	carbohydrate isomerase, AraD/FucA family
ORF02020	SAG1811	287	hexulose-6-phosphate isomerase, putative
ORF02021	SAG1812	221	hexulose-6-phosphate synthase, putative
ORF02022	SAG1813	161	PTS system, IIA component
ORF02023	SAG1814	92	PTS system, IIB component
ORF02024	SAG1815	479	transport protein SgaT, putative
ORF02025	SAG1816	205	hypothetical protein
ORF02026	SAG1817	157	hypothetical protein
ORF02027	SAG1818	430	adenylosuccinate synthetase
ORF02028	SAG1819	340	perfringolysin O regulator protein
ORF02029	SAG1820	224	conserved hypothetical protein
ORF02030	SAG1821	750	glutamatecysteine ligase-related protein
ORF02031	SAG1822	272	conserved hypothetical protein
ORF02032 ORF02033	SAG1823	418	conserved hypothetical protein
ORF02033	SAG1824	291	chaperonin, 33 kDa
ORF02034	SAG1825	325	NifR3/Smm1 family protein
ORF02036	SAG1826	213	deoxynucleoside kinase family protein
ORF02037	SAG1827	163	phosphinothricin N-acetyltransferase
OK 02037	SAG1828	815	ATP-dependent Clp protease, ATP-binding subunit
ORF02038	SAG1829	154	transcriptional regulator CtsR
ORF02039	SAG1830	153	conserved hypothetical protein
ORF02040	SAG1831	346	translation elongation factor Ts
ORF02041	SAG1832	256	ribosomal protein S2
ORF02042	SAG1833	186	alkyl hydroperoxide reductase, subunit C
ORF02043	SAG1834	510	alkyl hydroperoxide reductase, subunit F
ORF02044	SAG1835	134	conserved hypothetical protein
ORF02045	SAG1836	61	conserved hypothetical protein
ORF02046	SAG1837	468	lysin, putative
ORF02047	SAG1838	109	holin, putative
ORF02048	SAG1839	136	conserved hypothetical protein
ORF02049	SAG1840	112	hypothetical protein
ORF02050	SAG1841	76	conserved domain protein
ORF02051	SAG1842	1224	PbIB, putative
ORF02053	SAG1843	240	conserved hypothetical protein
ORF020~6	SAG1844	911	conserved hypothetical protein
ORF02057	SAG1845	42	hypothetical protein
ORF02058	SAG1846	158	hypothetical protein
ORF02059	SAG1847	227	conserved hypothetical protein
ORF02060	SAG1848	114	conserved hypothetical protein
ORF02061	SAG1849	115	hypothetical protein
			···/b



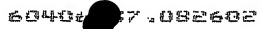
# Table 32: C nversion fORFR fN s. with SAG Ref Nos.

ORF R f No.	SAGxxxx Ref No.	aa	Ann tation
ORF02062	SAG1850	101	hypothetical protein
ORF02063	SAG1851 .	111	conserved domain protein
ORF02064	SAG1852	420	conserved domain protein
ORF02066	SAG1853	180	protease, putative
ORF02067	SAG1854	380	conserved hypothetical protein
ORF02068	SAG1855	570	terminase large subunit, putative
ORF02069	SAG1856	161	hypothetical protein
ORF02070	SAG1858	95	hypothetical protein
ORF02071	SAG1859	180	
011102077	UAG 1039	180	site-specific recombinase, phage integrase family
ORF02072	SAG1860	154	conserved hypothetical protein
ORF02073	SAG1861	119	transcriptional regulator, Cro/Cl family
ORF02075	SAG1862	86	hypothetical protein
ORF02076	SAG1863	138	single-strand binding protein
ORF02077	SAG1864	68	hypothetical protein
ORF02078	SAG1865	74	conserved hypothetical protein
ORF02079	SAG1866	109	conserved hypothetical protein
ORF02080	SAG1867	163	conserved hypothetical protein
ORF02081	SAG1868	134	hypothetical protein
ORF02082	SAG1869	437	type II DNA modification methyltransferase, putative
ORF02083	SAG1870	273	DNA replication protein DnaC, putative
ORF02084	SAG1871	248	conserved hypothetical protein
ORF02085	SAG1872	200	hypothetical protein
ORF02086	SAG1873	443	replicative DNA helicase
ORF02087	SAG1874	87	hypothetical protein
ORF02088	SAG1875	94	conserved hypothetical protein
ORF02089	SAG1876	176	HNH endonuclease family protein
ORF02090	SAG1877	236	antirepressor protein, putative
ORF02091	SAG1878	102	conserved domain protein
ORF02092	SAG1879	156	hypothetical protein
ORF02093	SAG1880	54	hypothetical protein
ORF02094	SAG1881	51	hypothetical protein
ORF02095	SAG1882	120	repressor protein, putative
ORF02097	SAG1884	134	hypothetical protein
ORF02098	SAG1885	356	site-specific recombinase, phage integrase family
	0.10.000	000	Site-specific recombinase, priage integrase family
ORF02100	SAG1886	32	hypothetical protein
ORF02101	SAG1887	689	Na+/H+ exchanger family protein
ORF02102	SAG1888	78	hypothetical protein
ORF02103	SAG1889	317	microcin immunity protein MccF, putative
ORF02104	SAG1890	631	endopeptidase O
ORF02105	SAG1891	327	oxidoreductase, Gfo/Idh/MocA family
ORF02107	SAG1892	358	membrane protein, putative
ORF02108	SAG1893	59	hypothetical protein
ORF02109	SAG1894	214	Cyclic nucleotide-binding domain protein
ORF02110	SAG1895	204	polypeptide deformylase
ORF02111	SAG1896	333	sugar binding transcriptional regulator RegR
ORF02112	SAG1897	634	conserved hypothetical protein
ORF02113	SAG1898	271	PTS system, IID component
ORF02114	SAG1899	288	PTS system, IIC component
ORF02115	SAG1900	164	PTS system, IIB component
ORF02116	SAG1901	398	glucuronyi hydrolase
ORF02118	SAG1902	144	
ORF02119	SAG1902		PTS system, IIA component
	1 0401903	34	hypothetical protein



# Tabl 32: Conv rsion of ORF R f Nos. with SAG Ref Nos.

ORF Ref No.	SAGxxxx Ref No.	aa	Ann tation
ORF02120	SAG1904	270	oxidoreductase, short-chain dehydrogenase/reductas
			<u>tamily</u>
ORF02121	SAG1905	212	conserved hypothetical protein
ORF02122	SAG1906	335	carbohydrate kinase, PfkB family
ORF02123	SAG1907	212	2-dehydro-3-deoxyphosphogluconate aldolase/4-
ODE02424	0404000	-:20	hydroxy-2-oxoglutarate aldolase
ORF02124	SAG1908	499	hypothetical protein
ORF02125 ORF02126	SAG1909	204	nitroreductase family protein
ORF02126	SAG1910 SAG1911	141	transcriptional regulator, MarR family
		1468	DNA polymerase III, alpha subunit, Gram-positive type
ORF02128	SAG1912	194	N-acetylmuramoyl-L-alanine amidase, family 4 protein
ORF02129	SAG1913	617	prolyl-tRNA synthetase
ORF02130	SAG1914	419	membrane-associated zinc metalloprotease, putative
			Parameter Emb metamoprotease, putante
ORF02131	SAG1915	264	phosphatidate cytidylyltransferase
ORF02132	SAG1916	250	undecaprenyl diphosphate synthase
ORF02133	SAG1917	113	preprotein translocase, YajC subunit
ORF02134	SAG1918	114	conserved hypothetical protein
ORF02135	SAG1919	387	malate oxidoreductase
ORF02136	SAG1920	445	citrate carrier protein, CCS family
ORF02137	SAG1921	508	sensor histidine kinase family protein
ORF02138	SAG1922	229	response regulator
ORF02139	SAG1923	331	UDP-glucose 4-epimerase
ORF02140	SAG1924	535	glucan 1,6-alpha-glucosidase
ORF02141	SAG1925	377	sugar ABC transporter, ATP-binding protein
ORF02142	SAG1926	283	helix-turn-helix domain protein, fis-type
ORF02143	SAG1927	298	lacX protein
ORF02144	SAG1928	325	tagatose 1,6-diphosphate aldolase
ORF02145	SAG1929	310	tagatose-6-phosphate kinase
ORF02146	SAG1930	171	galactose-6-phosphate isomerase, LacB subunit
ORF02147	SAG1931	141	galactose-6-phosphate isomerase, LacA subunit
ORF02148	. SAG1932	816	neuraminidase
ORF02149	SAG1933	482	PTS system, IIC component, putative
ORF02150	SAG1934	101	PTS system, IIB component, putative
ORF02152	SAG1935	157	PTS system, IIA component, putative
ORF02153	SAG1936	258	lactose phosphotransferase system repressor
ORF02156	SAG1937		streptococcal histidine triad family protein, degenerate FRAMESHIFT
ORF02157	SAG1938	307	adhesion lipoprotein, putative
ORF02158	SAG1939	147	conserved hypothetical protein TIGR00256
ORF02159	SAG1940	738	GTP pyrophosphokinase
ORF02160	SAG1941	800	2',3'-cyclic-nucleotide 2'-phosphodiesterase
ORF02161	SAG1942	151	nrdl protein, putative
ORF02162	SAG1943	345	conserved hypothetical protein
ORF02163	SAG1944	165	conserved hypothetical protein
ORF02164	SAG1945	345	iron ABC transporter, iron-binding protein
ORF02165	SAG1946	257	DNA-binding response regulator
ORF02166	SAG1947	549	conserved hypothetical protein
ORF02167	SAG1948	275	PTS system, IID component
ORF02168	SAG1949	269	PTS system, IIC component
ORF02169	SAG1950	163	PTS system, IIB component
ORF02170	SAG1951	141	PTS system, IIA component, putative
ORF02171	SAG1952	353	membrane protein, putative
ORF02172	SAG1953	60	hypothetical protein



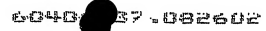
## Table 32: Conv rsi n of ORF Ref Nos. with SAG R f Nos.

ORF Ref No.	SAGxxxx Ref No.	aa	Ann tation
ORF02173	SAG1954	384	hypothetical protein
ORF02174	SAG1955	282	ABC transporter, ATP-binding protein
ORF02175	SAG1956	96	conserved domain protein
ORF02176	SAG1957	250	response regulator
ORF02177	SAG1958	276	conserved hypothetical protein
ORF02178	SAG1959	727	PTS system, IIABC components
ORF02179	SAG1960	551	
ORF02180	SAG1961		sensor histidine kinase
ORF02181	SAG1962	225	phosphate regulon response regulator PhoB
0/(10210)	SAG1902	218	phosphate transport system regulatory protein PhoU, putative
ORF02182	SAG1963	253	phosphate ABC transporter, ATP-binding protein
ORF02183	SAG1964	292	phosphate ABC transporter, permease protein
ORF02184	SAG1965	281	phosphate ABC transporter, permease protein
ORF02186	SAG1966	293	hemolysin precursor, putative
ORF02187	SAG1967	195	hypothetical protein
ORF02188	SAG1968	246	
ORF02189	SAG1969	317	conserved hypothetical protein TIGR00046
ORF02190	SAG1970	102	ribosomal protein L11 methyltransferase
ORF02191	SAG1970		conserved hypothetical protein
ORF02192		41	hypothetical protein
ORF02192	SAG1972	238	transcriptional regulator, MerR family
	SAG1973	156	acetyltransferase, GNAT family
ORF02195	SAG1974	152	MutT/nudix family protein
ORF02196	SAG1975	47	hypothetical protein
ORF02197	SAG1976	156	conserved hypothetical protein
ORF02198	SAG1977	163	acetyltransferase, GNAT family
ORF02199	SAG1978	422	ATPase, AAA family
ORF02201	SAG1979	253	hypothetical protein
ORF02202	SAG1980	300	ABC transporter, ATP-binding protein
ORF02203	SAG1981	68	hypothetical protein
ORF02205	SAG1982	359	transcriptional regulator, Cro/CI family
ORF02206	SAG1983	105	conserved hypothetical protein
ORF02207	SAG1984	188	conserved hypothetical protein TIGR00730
ORF02208	SAG1985	51	hypothetical protein
ORF02209	SAG1986	375	integrase, phage family, putative
ORF02210	SAG1987	61	conserved hypothetical protein
ORF02211	SAG1988	342	conserved hypothetical protein
ORF02212	SAG1989	139	hypothetical protein
ORF02213	SAG1990	127	hypothetical protein
ORF02214	SAG1991	204	transcriptional regulator, Cro/Cl family
ORF02215	SAG1992	518	conserved hypothetical protein
ORF02216	SAG1993	373	site-specific recombinase, phage integrase family
ORF02217	SAG1994	108	conserved hypothetical protein
ORF02219	SAG1995	210	hypothetical protein
ORF02221	SAG1996	263	cell wall anchor protein-related protein
ORF02223	SAG1998	182	hypothetical protein
ORF02224	SAG1997 SAG1998	457	
ORF02225	SAG1998 SAG1999		hypothetical protein
ORF02226		47	hypothetical protein
ORF02227	SAG2000	666	membrane protein, putative
ORF02228	SAG2001	756	conjugal transfer protein, interruption-C
ORF02228	SAG2002	129	IS1381, transposase OrfB
	SAG2003	127	IS1381, transposase OrfA
ORF02230	SAG2005	136	conserved hypothetical protein
ORF02231	SAG2006	88	conserved hypothetical protein
ORF02232	SAG2007	317	conserved hypothetical protein



# Table 32: Conversion f ORF R f Nos. with SAG Ref Nos.

ORF Ref No.	SAGxxxx Ref No.	aa	Ann tation
ORF02233	SAG2008	84	conserved hypothetical protein
ORF02234	SAG2009	88	conserved hypothetical protein
ORF02235	SAG2010	157	hypothetical protein
ORF02236	SAG2011	160	conserved hypothetical protein
ORF02237	SAG2012	90	hypothetical protein
ORF02238	SAG2013	189	hypothetical protein
ORF02239	SAG2014	449	hypothetical protein
ORF02240	SAG2015	99	transcriptional regulator, Cro/CI family
ORF02241	SAG2016	125	hypothetical protein
ORF02242	SAG2017	429	transcriptional regulator, Cro/CI family
ORF02243	SAG2018	553	FtsK/SpollIE family protein
ORF02244	SAG2019	153	hypothetical protein
ORF02245	SAG2020	98	hypothetical protein
ORF02246	SAG2021	826	cell wall surface anchor family protein
ORF02247	SAG2022	417	transposase, ISL3 family
ORF02249	SAG2023	546	mercuric reductase
ORF02250	SAG2024	130	
	3.102027	,,,,	mercuric resistance operon regulatory protein MerR
ORF02251	-SAG2025	522	Mn2+/Fe2+ transporter, NRAMP family
ORF02252	SAG2026	240	membrane protein, putative
ORF02253	SAG2027	205	ABC transporter, ATP-binding protein
ORF02254	SAG2028	36	conserved hypothetical protein
ORF02255	SAG2029	284	streptomycin resistance protein
ORF02257	SAG2030	130	hypothetical protein
ORF02258	SAG2031	202	hypothetical protein
ORF02259	SAG2032	111	conserved hypothetical protein
ORF02260	SAG2033	162	acetyltransferase, GNAT family
ORF02261	SAG2034	247	membrane protein, putative
ORF02262	SAG2035	300	ABC transporter, ATP-binding protein
ORF02263	SAG2036	68	hypothetical protein
ORF02264	SAG2037	358	transcriptional regulator, Cro/CI family
ORF02265	SAG2038	204	PAP2 family protein
ORF02266	SAG2039	98	conserved hypothetical protein
ORF02267	SAG2040	186	conserved hypothetical protein TIGR00730
ORF02268	SAG2041	287	protease, putative
ORF02269	SAG2042	100	rhodanese family protein
ORF02270	SAG2043	255	cAMP factor
ORF02271	SAG2044	62	hypothetical protein
ORF02272	SAG2045	179	DNA topology modulation protein FlaR, putative
ORF02273	SAG2046	361	glycerol dehydrogenase, putative
ORF02274	SAG2047	235	conserved hypothetical protein
ORF02275	SAG2048	614	5-methyltetrahydrofolatehomocysteine
			methyltransferase, putative
ORF02276	SAG2049	745	5-methyltetrahydropteroyltriglutamatehomocysteine
OPENNIZZ	0400000		methyltransferase
ORF02277 ORF02278	SAG2050	107	conserved hypothetical protein
URF022/8	SAG2051	230	branched-chain amino acid transport protein AzlC,
ORF02279	SAG2052	44	putative
ORF02280	SAG2052 SAG2053	41	hypothetical protein
ORF02281	SAG2053 SAG2054	1570	serine protease, subtilase family, putative
ORF02282	SAG2054 SAG2055	228	DNA-binding response regulator
ORF02283	SAG2055 SAG2056	462	sensor histidine kinase
ORF02285	SAG2056 SAG2057	202	chromosome assembly-related protein
ORF02286	SAG2057 SAG2058	833	leucyl-tRNA synthetase
	UNG2036	415	major facilitator family protein



# Tabl 32: Conversi n f ORF R f Nos. with SAG Ref N s.

ORF02287         SAG2059         281         conserved hypothetical protein           ORF02288         SAG2080         398         glycosyl transferase, family 8           ORF02289         SAG2081         401         glycosyl transferase, family 8           ORF02291         SAG2062         179         transcription antitermination protein NusG           ORF02292         SAG2064         57         preprotein translocase, Sect subunit, putative           ORF02293         SAG2068         57         preprotein translocase, Sect subunit, putative           ORF02294         SAG2068         546         Lyme disease proteins of unknown function, putative           ORF02295         SAG2068         546         Lyme disease proteins of unknown function, putative           ORF02296         SAG2068         546         Lyme disease proteins of unknown function, putative           ORF02297         SAG2070         223         deoxyribose-phosphate aldolase           ORF02298         SAG2071         223         deoxyribose-phosphate aldolase           ORF02299         SAG2071         223         deoxyribose-phosphate aldolase           ORF02290         SAG2071         240         Na respective phosphate aldolase           ORF02300         SAG2071         240         Na respective phosphate aldolase	ORF Ref No.	SAGxxxx Ref N .	aa	Annotation
ORF02288         SAG2060         398         glycosyl transferase, family 8           ORF02289         SAG2061         401         glycosyl transferase, family 8           ORF02290         SAG2062         179         branchina mitlermination protein NusG           ORF02291         SAG2084         57         perprotein translocase, SecE subunit, putative           ORF02293         SAG2066         773         periotilith-inhiding protein 2A           ORF02294         SAG2067         294         blosomal large subunit pseudouridine synthase, RluD subramily           ORF02295         SAG2068         546         Lyme disease proteins of unknown function, putative subramily           ORF02296         SAG2069         403         phosphopentomutase           ORF02297         SAG2070         223         deoxyribose-phosphate aldolase           ORF02298         SAG2071         400         Na+ dependent nucleoside transporter           ORF02301         SAG2072         259         unitide phosphorylase           ORF02301         SAG2073         245         transcriptional regulator, GnIR family           ORF02302         SAG2074         540         80 kda chaperonin, 10 kDa           ORF02303         SAG2075         267         ABC transporter, ATP-binding protein <t< td=""><td>ORF02287</td><td>SAG2059</td><td></td><td></td></t<>	ORF02287	SAG2059		
ORF02289         SAG2061         401         dycosyl transferase, family 8           ORF02291         SAG2062         178         transcription antifermination protein NusG           ORF02292         SAG2063         530         pathogenicity protein, putative           ORF02293         SAG2066         773         pericillin-briding protein 2A           ORF02294         SAG2067         294         ribosomal large subunit pseudouridine synthase, RtuD subfamily           ORF02295         SAG2068         546         Lyme disease proteins of unknown function, putative           ORF02296         SAG2069         403         phosphopentomutase           ORF02297         SAG2070         223         deoxyribose-phosphate aldolase           ORF02298         SAG2071         400         Na4-dependent nuclooside transporter           ORF02290         SAG2072         259         uridine phosphorylase           ORF02301         SAG2073         245         transcriptional regulator, GriR family           ORF02302         SAG2074         540         60 kda chaperonin           ORF02303         SAG2076         227         ABC transporter, Permease protein           ORF02304         SAG2077         296         ABC transporter, permease protein           ORF02305         S	ORF02288	SAG2060	398	
ORF02290         SAG2062         179         transcription antifermination protein NusG           ORF02291         SAG2064         67         preprotein translocase, SecE subunit, putative           ORF02292         SAG2064         67         preprotein translocase, SecE subunit, putative           ORF02293         SAG2066         773         perincillin-binding protein 2A           ORF02294         SAG2067         294         biosomal large subunit pseudouridine synthase, RtuD subfamily           ORF02295         SAG2068         546         Lyme disease proteins of unknown function, putative subfamily           ORF02296         SAG2069         403         phosphopentomutase           ORF02297         SAG2070         223         deoxyribose-phosphate aldolase           ORF02298         SAG2071         400         Na+ dependent nucleoside transporter           ORF02300         SAG2073         245         transcriptional regulator, GnIR family           ORF02301         SAG2073         245         transcriptional regulator, GnIR family           ORF02302         SAG2076         94         chaperonin, 10 kDa           ORF02303         SAG2077         298         ABC transporter, ATP-binding protein           ORF02304         SAG2078         320         lipoprotein, putative	ORF02289	SAG2061	401	
ORF02291         SAG2084         630         pathogenicity protein, putative           ORF02292         SAG2066         773         perpotein translocase, SecE subunit, putative           ORF02294         SAG2066         773         perpotein translocase, SecE subunit, putative           ORF02295         SAG2068         294         ribosomal large subunit pseudouridine synthase, RiuD subdamily           ORF02296         SAG2069         403         phosphopentomutase           ORF02297         SAG2070         223         deoxyribose-phosphate aidolase           ORF02298         SAG2071         400         Na+ dependent nucleoside transporter           ORF02290         SAG2072         259         uridine phosphorylase           ORF02301         SAG2077         245         transcriptional regulator, OntR family           ORF02302         SAG2074         540         60 kda chaperonin           ORF02303         SAG2075         297         ABC transporter, ATP-binding protein           ORF02304         SAG2077         299         ABC transporter, Permease protein           ORF02305         SAG2078         320         lipoprotein, putative           ORF02310         SAG2083         285         hydrolase, haloacid dehalogenase-like family           ORF02311	ORF02290	SAG2062	179	
ORF02292         SAG2084         57         preprotein translocase, SecE subunii, putative           ORF02294         SAG2067         294         inbosomal large subunit pseudouridine synthase, RluD subfamily           ORF02295         SAG2068         546         Lyme disease proteins of unknown function, putative           ORF02296         SAG2069         403         phosphopentomutase           ORF02297         SAG2070         223         deoxyribose-phosphate aldolase           ORF02298         SAG2071         400         Na+ dependent nucloside transporter           ORF02300         SAG2072         259         uridine phosphorylase           ORF02301         SAG2073         245         transcriptional regulator, GriR family           ORF02302         SAG2073         245         transcriptional regulator, GriR family           ORF02303         SAG2076         247         transcriptional regulator, GriR family           ORF02303         SAG2077         298         ABC transporter, ATP-binding protein           ORF02303         SAG2078         320         lipoprotein, putative           ORF02304         SAG2078         320         lipoprotein, putative           ORF02305         SAG2080         286         glyoxalase family protein           ORF02310	ORF02291	SAG2063	630	
ORF02293         SAG2066         773         penicilin-binding protein ZA           ORF02294         SAG2067         294         ribosomal large subuniti pseudouridine synthase, RluD subfamily           ORF02295         SAG2068         546         Lyme disease proteins of unknown function, putative           ORF02296         SAG2069         403         phosphopentormutase           ORF02297         SAG2079         223         deoxyribose-phosphate aldolase           ORF02298         SAG2071         400         Na+ dependent nucleoside transporter           ORF02300         SAG2072         259         unidine phosphorylase           ORF02301         SAG2073         245         transcriptional regulator, GriRR family           ORF02302         SAG2074         540         80 kda chaperonin           ORF02303         SAG2076         291         ABC transporter, Permease protein           ORF02304         SAG2077         298         ABC transporter, permease protein           ORF02305         SAG2078         2265         hydrolase, haloacid dehalogenase-like family           ORF02306         SAG2081         243         conserved hypothetical protein           ORF02310         SAG2083         163         acetytransferase, GNAT family           ORF02311         <	ORF02292	SAG2064	57	
ORF02294         SAG2087         294         Inbosomal large subunit pseudouridine synthase, RtluD subfamily           ORF02295         SAG2088         546         Lyme disease proteins of unknown function, putative           ORF02296         SAG2089         403         phosphopentomutase           ORF02297         SAG2070         223         deoxyrilose-phosphate aldolase           ORF02300         SAG2071         400         Na+ dependent nucleoside transporter           ORF02301         SAG2073         245         transcriptional regulator, GriR family           ORF02302         SAG2075         245         transcriptional regulator, GriR family           ORF02303         SAG2076         259         al Caransporter, ATP-binding protein           ORF02303         SAG2077         298         ABC transporter, Permease protein           ORF02305         SAG2078         320         lipoprotein, putative           ORF02307         SAG2078         320         lipoprotein, putative           ORF02308         SAG2079         285         hydrolase, haloacid dehalogenase-like family           ORF02309         SAG2081         243         conserved hypothetical protein           ORF02310         SAG2081         243         conserved hypothetical protein           ORF02312	ORF02293	SAG2066	773	
Subfamily	ORF02294	SAG2067	294	
ORF02298				subfamily
ORF02297         SAG2070         223         deoxyribose-phosphate aldolase           ORF02298         SAG2071         400         Na+ dependent nucleoside transporter           ORF02300         SAG2072         259         uridine phosphorylase           ORF02301         SAG2073         245         transcriptional regulator, GnIR family           ORF02302         SAG2076         540         60 kda chaperonin           ORF02303         SAG2076         94         chaperonin, 10 kDa           ORF02305         SAG2076         267         ABC transporter, ATP-binding protein           ORF02306         SAG2078         280         lipoprotein, putative           ORF02307         SAG2080         286         glyoxalase family protein           ORF02308         SAG2078         230         lipoprotein, putative           ORF02309         SAG2080         286         glyoxalase family protein           ORF02310         SAG2080         286         glyoxalase family protein           ORF02311         SAG2082         205         anaerobic ribonucleoside-triphosphate reductase           ORF02312         SAG2083         163         acetyltransferase, GNAT family           ORF02313         SAG2084         47         conserved hypothetical protein		SAG2068	546	Lyme disease proteins of unknown function, putative
ORF02297         SAG2070         223         deoxyribose-phosphate aldolase           ORF02300         SAG2071         400         Na+ dependent nucleoside transporter           ORF02301         SAG2073         245         uridine phosphorylase           ORF02301         SAG2073         245         transcriptional regulator, GntR family           ORF02302         SAG2074         540         60 kda chaperonin           ORF02305         SAG2075         94         chaperonin, 10 kDa           ORF02306         SAG2076         267         ABC transporter, ATP-binding protein           ORF02306         SAG2077         298         ABC transporter, permease protein           ORF02307         SAG2078         320         lipoprotein, putative           ORF02308         SAG2079         265         hydrolase family protein           ORF02309         SAG2080         286         alyoxalase family protein           ORF02310         SAG2081         243         conserved hypothetical protein           ORF02311         SAG2082         205         anaerobic ribonucleoside-triphosphate reductase           ORF02312         SAG2083         310         virulence factor Mvlm, putative           ORF02313         SAG2084         310         virulence factor Mv	ORF02296	SAG2069	403	phosphopentomutase
ORF02298         SAG2071         400         Na+ dependent nucleoside transporter           ORF02300         SAG2072         259         uridine phosphorylase           ORF02301         SAG2073         245         franscriptional regulator, GritR family           ORF02302         SAG2074         540         80 kda chaperonin           ORF02303         SAG2076         267         ABC transporter, ATP-binding protein           ORF02306         SAG2077         298         ABC transporter, permease protein           ORF02307         SAG2078         320         lipoprotein, putative           ORF02308         SAG2079         285         hydrolase, haloacid dehalogenase-like family           ORF02309         SAG2080         286         glyoxalase family protein           ORF02310         SAG2081         243         conserved hypothetical protein           ORF02311         SAG2082         205         anaerobic ribonucleoside-triphosphate reductase activating protein           ORF02312         SAG2083         183         acelytransferase, GNAT family           ORF02313         SAG2084         310         virulence factor MvIM, putative           ORF02314         SAG2085         47         conserved hypothetical protein           ORF02315         SAG2086	ORF02297	SAG2070	223	
ORF023001         SAG2072         259         uridine phosphorylase           ORF02301         SAG2073         245         transcriptional regulator, GntR family           ORF02302         SAG2074         540         60 kda chaperonin           ORF02303         SAG2076         94         chaperonin, 10 kDa           ORF02306         SAG2077         288         ABC transporter, ATP-binding protein           ORF02307         SAG2078         320         lipoprotein, putative           ORF02308         SAG2079         285         hydrolase, haloacid dehalogenase-like family           ORF02309         SAG2080         286         glyoxalase family protein           ORF02310         SAG2081         243         conserved hypothetical protein           ORF02311         SAG2082         205         anaerobic ribonucleoside-triphosphate reductase activating protein           ORF02312         SAG2083         163         acetyltransferase, GNAT family           ORF02313         SAG2084         310         virulence factor MvIM, putative           ORF02314         SAG2084         310         virulence factor MvIM, putative           ORF02315         SAG2086         723         anaerobic ribonucleoside-triphosphate reductase           ORF02316         SAG2087		SAG2071	400	
ORF02301         SAG2073         245         transcriptional regulator, GntR family           ORF02303         SAG2074         540         60 kda chaperonin           ORF02303         SAG2076         94         chaperonin, 10 kDa           ORF02305         SAG2076         267         ABC transporter, ATP-binding protein           ORF02306         SAG2077         298         ABC transporter, Permease protein           ORF02307         SAG2078         320         lipoprotein, putative           ORF02308         SAG2079         265         hydrolase, haloacid dehalogenase-like family           ORF02309         SAG2080         286         glyoxalase family protein           ORF02310         SAG2081         243         conserved hypothetical protein           ORF02311         SAG2082         205         anaerobic ribonucleoside-triphosphate reductase activating protein           ORF02312         SAG2083         163         acetyltransferase, GNAT family           ORF02313         SAG2084         310         virulence factor MviM, putative           ORF02314         SAG2085         47         conserved hypothetical protein           ORF02315         SAG2086         723         anaerobic ribonucleoside-triphosphate reductase           ORF02316         SAG2087	ORF02300	SAG2072	259	
ORF02302         SAG2074         540         60 kda chaperonin           ORF02303         SAG2075         94         chaperonin, 10 kDa           ORF02305         SAG2076         267         ABC transporter, ATP-binding protein           ORF02306         SAG2077         298         ABC transporter, permease protein           ORF02307         SAG2079         265         hydrolase, haloacid dehalogenase-like family           ORF02309         SAG2080         286         glyoxalase family protein           ORF02310         SAG2081         243         conserved hypothetical protein           ORF02311         SAG2082         205         anaerobic ribonucleoside-triphosphate reductase activating protein           ORF02312         SAG2083         163         acetyltransferase, GNAT family           ORF02313         SAG2084         310         virulence factor whilk, putative           ORF02314         SAG2085         47         conserved hypothetical protein           ORF02315         SAG2086         723         anaerobic ribonucleoside-triphosphate reductase           ORF02316         SAG2087         495         conserved hypothetical protein           ORF02317         SAG2088         40         hypothetical protein           ORF02318         SAG2098	ORF02301	SAG2073	245	
ORF02305   SAG2076   267   ABC transporter, ATP-binding protein   ORF02306   SAG2077   298   ABC transporter, permease protein   ORF02307   SAG2078   320   Ilipoprotein, putative   ORF02308   SAG2079   265   hydrolase, haloacid dehalogenase-like family   ORF02309   SAG2080   286   glyoxalase family protein   ORF02310   SAG2081   243   conserved hypothetical protein   ORF02311   SAG2082   205   anaerobic ribonucleoside-triphosphate reductase activating protein   ORF02312   SAG2083   163   acetyltransferase, GNAT family   ORF02313   SAG2084   310   virulence factor MvliM, putative   ORF02314   SAG2085   47   conserved hypothetical protein   ORF02315   SAG2086   723   anaerobic ribonucleoside-triphosphate reductase   ORF02316   SAG2087   495   conserved hypothetical protein   ORF02317   SAG2088   40   hypothetical protein   ORF02319   SAG2089   105   conserved hypothetical protein   ORF02319   SAG2090   136   conserved hypothetical protein   ORF02320   SAG2091   88   conserved hypothetical protein   ORF02321   SAG2092   132   conserved hypothetical protein   ORF02322   SAG2093   379   recA protein   ORF02323   SAG2094   conserved hypothetical protein   ORF02325   SAG2095   183   DNA-3-methyladenine glycosylase     ORF02326   SAG2097   418   transporter, putative   ORF02330   SAG2099   33   hypothetical protein   ORF02329   SAG2099   418   transporter, putative   ORF02330   SAG2099   33   hypothetical protein   ORF02331   SAG2100   67   cold shock protein, CSD family   ORF02332   SAG2101   858   DNA mismatch repair protein HexA   ORF02333   SAG2104   102   conserved hypothetical protein   ORF02334   SAG2105   290   conserved hypothetical protein   ORF02337   SAG2106   364   conserved hypothetical protein   ORF02338   SAG2104   102   conserved hypothetical protein   ORF02339   SAG2104   588   DNA mismatch repair protein HexA   ORF02331   SAG2105   290   conserved hypothetical protein   ORF02333   SAG2104   102   conserved hypothetical protein   ORF02339   SAG2105   390   conserved hypothetical protein			540	
ORF02305         SAG2076         267         ABC transporter, ATP-binding protein           ORF02306         SAG2077         298         ABC transporter, permease protein           ORF02308         SAG2078         320         lipoprotein, putative           ORF02308         SAG2080         286         hydrolase, haloacid dehalogenase-like family           ORF02309         SAG2081         243         conserved hypothetical protein           ORF02311         SAG2082         205         anaerobic ribonucleoside-triphosphate reductase activating protein           ORF02312         SAG2083         163         acetyltransferase, GNAT family           ORF02313         SAG2084         310         virulence factor M/IM, putative           ORF02314         SAG2085         47         conserved hypothetical protein           ORF02315         SAG2085         47         conserved hypothetical protein           ORF02316         SAG2087         495         conserved hypothetical protein           ORF02318         SAG2089         105         conserved hypothetical protein           ORF02319         SAG2098         105         conserved hypothetical protein           ORF02321         SAG2091         186         conserved hypothetical protein           ORF02322         SA		SAG2075	94	chaperonin, 10 kDa
ORF02306         SAG2077         298         ABC transporter, permease protein           ORF02307         SAG2078         320         lipoprotein, putative           ORF02308         SAG2079         265         hydrolase, haloacid dehalogenase-like family           ORF02309         SAG2080         286         glyoxalase family protein           ORF02310         SAG2081         243         conserved hypothetical protein           ORF02311         SAG2082         205         anaerobic ribonucleoside-triphosphate reductase activating protein           ORF02312         SAG2083         163         acetyltransferase, GNAT family           ORF02313         SAG2084         310         virulence factor MvIM, putative           ORF02314         SAG2085         47         conserved hypothetical protein           ORF02315         SAG2086         723         anaerobic ribonucleoside-triphosphate reductase           ORF02316         SAG2087         495         conserved hypothetical protein           ORF02317         SAG2088         40         hypothetical protein           ORF02318         SAG2089         105         conserved hypothetical protein           ORF02329         SAG2091         38         conserved hypothetical protein           ORF02321         SAG2092	ORF02305	SAG2076	267	
ORF02307         SAG2078         320         lipoprotein, putative           ORF02308         SAG2079         285         hydrolase, haloacid dehalogenase-like family           ORF02309         SAG2080         286         glyoxalase family protein           ORF02310         SAG2081         243         conserved hypothetical protein           ORF02311         SAG2082         205         anaerobic ribonucleoside-triphosphate reductase activating protein           ORF02312         SAG2083         163         acetyltransferase, GNAT family           ORF02313         SAG2084         310         virulence factor MvIM, putative           ORF02314         SAG2085         47         conserved hypothetical protein           ORF02314         SAG2086         723         anaerobic ribonucleoside-triphosphate reductase           ORF02314         SAG2085         47         conserved hypothetical protein           ORF02316         SAG2087         495         conserved hypothetical protein           ORF02317         SAG2088         40         hypothetical protein           ORF02318         SAG2099         136         conserved hypothetical protein           ORF02319         SAG2091         38         conserved hypothetical protein           ORF02321         SAG2092 <td></td> <td>SAG2077</td> <td>298</td> <td>ABC transporter, permease protein</td>		SAG2077	298	ABC transporter, permease protein
ORF02308         SAG2079         265         hydrolase, haloacid dehalogenase-like family           ORF02309         SAG2081         286         glyoxalase family protein           ORF02310         SAG2081         243         conserved hypothetical protein           ORF02311         SAG2082         205         anaerobic ribonucleoside-triphosphate reductase activating protein           ORF02312         SAG2083         163         acetyltransferase, GNAT family           ORF02313         SAG2084         310         virulence factor MvIM, putative           ORF02314         SAG2085         47         conserved hypothetical protein           ORF02315         SAG2086         723         anaerobic ribonucleoside-triphosphate reductase           ORF02316         SAG2086         723         anaerobic ribonucleoside-triphosphate reductase           ORF02318         SAG2087         495         conserved hypothetical protein           ORF02317         SAG2088         40         hypothetical protein           ORF02318         SAG2089         105         conserved hypothetical protein           ORF02329         SAG2099         136         conserved hypothetical protein           ORF02320         SAG2091         88         conserved hypothetical protein           ORF02322 </td <td></td> <td>SAG2078</td> <td>320</td> <td>lipoprotein, putative</td>		SAG2078	320	lipoprotein, putative
ORF02309         SAG2080         286         glyoxalase family protein           ORF02310         SAG2081         243         conserved hypothetical protein           ORF02311         SAG2082         205         anaerobic ribonucleoside-triphosphate reductase activating protein           ORF02312         SAG2083         163         acetyltransferase, GNAT family           ORF02313         SAG2084         310         virulence factor MvIM, putative           ORF02314         SAG2085         47         conserved hypothetical protein           ORF02315         SAG2086         723         anaerobic ribonucleoside-triphosphate reductase           ORF02316         SAG2087         495         conserved hypothetical protein           ORF02317         SAG2088         40         hypothetical protein           ORF02318         SAG2089         105         conserved hypothetical protein           ORF02319         SAG2099         136         conserved hypothetical protein           ORF02319         SAG2090         136         conserved hypothetical protein           ORF02321         SAG2091         88         conserved hypothetical protein           ORF02322         SAG2093         379         recA protein           ORF02323         SAG2094         competen		SAG2079	265	
ORF02310         SAG2081         243         conserved hypothetical protein           ORF02311         SAG2082         205         anaerobic ribonucleoside-triphosphate reductase activating protein           ORF02312         SAG2083         163         acetyltransferase, GNAT family           ORF02313         SAG2084         310         virulence factor MvIM, putative           ORF02314         SAG2085         47         conserved hypothetical protein           ORF02315         SAG2086         723         anaerobic ribonucleoside-triphosphate reductase           ORF02316         SAG2087         495         conserved hypothetical protein           ORF02317         SAG2088         40         hypothetical protein           ORF02318         SAG2089         105         conserved hypothetical protein           ORF02319         SAG2099         136         conserved hypothetical protein           ORF02320         SAG2091         88         conserved hypothetical protein           ORF02321         SAG2092         132         conserved hypothetical protein           ORF02322         SAG2093         379         recA protein           ORF02323         SAG2094         competence/damage-inducible protein CinA           FRAMESHIFT         recappetence/damage-inducible protein		SAG2080	286	
ORF02311         SAG2082         205         anaerobic ribonucleoside-triphosphate reductase activating protein           ORF02312         SAG2083         163         acetyltransferase, GNAT family           ORF02313         SAG2084         310         virulence factor MvIM, putative           ORF02314         SAG2085         47         conserved hypothetical protein           ORF02315         SAG2086         723         anaerobic ribonucleoside-triphosphate reductase           ORF02316         SAG2087         495         conserved hypothetical protein           ORF02317         SAG2088         40         hypothetical protein           ORF02318         SAG2089         105         conserved hypothetical protein           ORF02319         SAG2090         136         conserved hypothetical protein TIGR00250           ORF02321         SAG2099         136         conserved hypothetical protein TIGR00250           ORF02322         SAG2091         38         conserved hypothetical protein           ORF02321         SAG2092         132         conserved hypothetical protein           ORF02322         SAG2093         379         recA protein           ORF02323         SAG2094         competence/damage-inducible protein CinA FRAMESHIFT           ORF02325         SAG2095<		SAG2081	243	
CRF02312   SAG2083   163   acetyltransferase, GNAT family	ORF02311	SAG2082	205	
ORF02313         SAG2084         310         virulence factor MWIM, putative           ORF02314         SAG2085         47         conserved hypothetical protein           ORF02315         SAG2086         723         anaerobic ribonucleoside-triphosphate reductase           ORF02316         SAG2087         495         conserved hypothetical protein           ORF02317         SAG2088         40         hypothetical protein           ORF02318         SAG2089         105         conserved hypothetical protein           ORF02319         SAG2090         136         conserved hypothetical protein           ORF02320         SAG2091         136         conserved hypothetical protein           ORF02321         SAG2092         132         conserved hypothetical protein           ORF02322         SAG2091         188         conserved hypothetical protein           ORF02323         SAG2092         132         conserved hypothetical protein           ORF02324         SAG2093         379         recA protein           ORF02325         SAG2094         competence/damage-inducible protein CinA           FRAMESHIFT         competence/damage-inducible protein CinA           ORF02327         SAG2096         196         Holliday junction DNA helicase RuvA				activating protein
ORF02314         SAG2085         47         conserved hypothetical protein           ORF02315         SAG2086         723         anaerobic ribonucleoside-triphosphate reductase           ORF02316         SAG2087         495         conserved hypothetical protein           ORF02317         SAG2088         40         hypothetical protein           ORF02318         SAG2089         105         conserved hypothetical protein           ORF02319         SAG2090         136         conserved hypothetical protein           ORF02320         SAG2091         88         conserved hypothetical protein           ORF02321         SAG2092         132         conserved hypothetical protein           ORF02322         SAG2093         379         recA protein           ORF02323         SAG2094         conserved hypothetical protein           ORF02323         SAG2094         conserved hypothetical protein           ORF02323         SAG2093         379         recA protein           ORF02323         SAG2094         conserved hypothetical protein           ORF02325         SAG2095         183         DNA-3-methyladenine glycosylase I           ORF02326         SAG2096         196         Holliday junction DNA helicase RuvA           ORF02327 <td< td=""><td></td><td></td><td>163</td><td></td></td<>			163	
ORF02315         SAG2086         723         anaerobic ribonucleoside-triphosphate reductase           ORF02316         SAG2087         495         conserved hypothetical protein           ORF02317         SAG2088         40         hypothetical protein           ORF02318         SAG2088         40         hypothetical protein           ORF02319         SAG2089         136         conserved hypothetical protein           ORF02320         SAG2091         88         conserved hypothetical protein           ORF02321         SAG2092         132         conserved hypothetical protein           ORF02322         SAG2093         379         recA protein           ORF02323         SAG2094         competence/damage-inducible protein CinA           FRAMESHIFT         competence/damage-inducible protein CinA           ORF02325         SAG2095         183         DNA-3-methyladenine glycosylase I           ORF02326         SAG2096         196         Holliday junction DNA helicase RuvA           ORF02327         SAG2098         659         DNA mismatch repair protein HexB           ORF02339         SAG2099         33         hypothetical protein           ORF02331         SAG2100         67         cold shock protein, CSD family           ORF02332<				
ORF02316 SAG2087 495 conserved hypothetical protein ORF02317 SAG2088 40 hypothetical protein ORF02318 SAG2089 105 conserved hypothetical protein ORF02319 SAG2090 136 conserved hypothetical protein TIGR00250 ORF02320 SAG2091 88 conserved hypothetical protein ORF02321 SAG2092 132 conserved hypothetical protein ORF02322 SAG2093 379 recA protein ORF02323 SAG2094 competence/damage-inducible protein CinA FRAMESHIFT ORF02325 SAG2095 183 DNA-3-methyladenine glycosylase I ORF02327 SAG2096 196 Holliday junction DNA helicase RuvA ORF02328 SAG2098 659 DNA mismatch repair protein HexB ORF02330 SAG2099 33 hypothetical protein ORF02331 SAG2100 67 cold shock protein, CSD family ORF02332 SAG2101 858 DNA mismatch repair protein HexA ORF02333 SAG2102 145 arginine repressor ArgR, putative ORF02334 SAG2103 563 arginyl-tRNA synthetase ORF02337 SAG2105 290 conserved hypothetical protein ORF02338 SAG2105 290 conserved hypothetical protein ORF02339 SAG2107 583 aspartyl-tRNA synthetase ORF02339 SAG2107 583 aspartyl-tRNA synthetase				
ORF02317         SAG2088         40         hypothetical protein           ORF02318         SAG2089         105         conserved hypothetical protein           ORF02319         SAG2090         136         conserved hypothetical protein           ORF02320         SAG2091         88         conserved hypothetical protein           ORF02321         SAG2092         132         conserved hypothetical protein           ORF02322         SAG2093         379         recA protein           ORF02323         SAG2094         competence/damage-inducible protein CinA FRAMESHIFT           ORF02325         SAG2095         183         DNA-3-methyladenine glycosylase I           ORF02327         SAG2096         196         Holliday junction DNA helicase RuvA           ORF02328         SAG2097         418         transporter, putative           ORF02329         SAG2098         659         DNA mismatch repair protein HexB           ORF02330         SAG2099         33         hypothetical protein           ORF02331         SAG2100         67         cold shock protein, CSD family           ORF02332         SAG2101         858         DNA mismatch repair protein HexA           ORF02333         SAG2102         145         arginnie repressor ArgR, putative     <		SAG2086	723	anaerobic ribonucleoside-triphosphate reductase
ORF02317         SAG2088         40         hypothetical protein           ORF02318         SAG2089         105         conserved hypothetical protein           ORF02319         SAG2090         136         conserved hypothetical protein TIGR00250           ORF02320         SAG2091         88         conserved hypothetical protein           ORF02321         SAG2092         132         conserved hypothetical protein           ORF02322         SAG2093         379         recA protein           ORF02323         SAG2094         competence/damage-inducible protein CinA FRAMESHIFT           ORF02325         SAG2094         competence/damage-inducible protein CinA FRAMESHIFT           ORF02327         SAG2095         183         DNA-3-methyladenine glycosylase I           ORF02328         SAG2096         196         Holliday junction DNA helicase RuvA           ORF02328         SAG2097         418         transporter, putative           ORF02330         SAG2098         659         DNA mismatch repair protein HexB           ORF02331         SAG2100         67         cold shock protein, CSD family           ORF02332         SAG2101         858         DNA mismatch repair protein HexA           ORF02333         SAG2102         145         arginyl-tRNA synthetase		SAG2087	495	conserved hypothetical protein
ORF02319         SAG2090         136         conserved hypothetical protein TIGR00250           ORF02320         SAG2091         88         conserved hypothetical protein           ORF02321         SAG2092         132         conserved hypothetical protein           ORF02322         SAG2093         379         recA protein           ORF02323         SAG2094         competence/damage-inducible protein CinA FRAMESHIFT           ORF02325         SAG2095         183         DNA-3-methyladenine glycosylase I           ORF02327         SAG2096         196         Holliday junction DNA helicase RuvA           ORF02328         SAG2097         418         transporter, putative           ORF02329         SAG2098         659         DNA mismatch repair protein HexB           ORF02330         SAG2099         33         hypothetical protein           ORF02331         SAG2100         67         cold shock protein, CSD family           ORF02332         SAG2101         858         DNA mismatch repair protein HexA           ORF02333         SAG2102         145         arginine repressor ArgR, putative           ORF02334         SAG2103         563         arginyl-tRNA synthetase           ORF02337         SAG2105         290         conserved hypothetical prot		SAG2088	40	
ORF02319         SAG2090         136         conserved hypothetical protein TIGR00250           ORF02320         SAG2091         88         conserved hypothetical protein           ORF02321         SAG2092         132         conserved hypothetical protein           ORF02322         SAG2093         379         recA protein           ORF02323         SAG2094         competence/damage-inducible protein CinA FRAMESHIFT           ORF02325         SAG2095         183         DNA-3-methyladenine glycosylase I           ORF02327         SAG2096         196         Holliday junction DNA helicase RuvA           ORF02328         SAG2097         418         transporter, putative           ORF02329         SAG2098         659         DNA mismatch repair protein HexB           ORF02330         SAG2099         33         hypothetical protein           ORF02331         SAG2100         67         cold shock protein, CSD family           ORF02332         SAG2101         858         DNA mismatch repair protein HexA           ORF02333         SAG2102         145         arginine repressor ArgR, putative           ORF02334         SAG2103         563         arginyl-tRNA synthetase           ORF02337         SAG2105         290         conserved hypothetical prot		SAG2089	105	
ORF02320         SAG2091         88         conserved hypothetical protein           ORF02321         SAG2092         132         conserved hypothetical protein           ORF02322         SAG2093         379         recA protein           ORF02323         SAG2094         competence/damage-inducible protein CinA FRAMESHIFT           ORF02325         SAG2095         183         DNA-3-methyladenine glycosylase I           ORF02327         SAG2096         196         Holliday junction DNA helicase RuvA           ORF02328         SAG2097         418         transporter, putative           ORF02329         SAG2098         659         DNA mismatch repair protein HexB           ORF02330         SAG2099         33         hypothetical protein           ORF02331         SAG2100         67         cold shock protein, CSD family           ORF02332         SAG2101         858         DNA mismatch repair protein HexA           ORF02333         SAG2102         145         arginine repressor ArgR, putative           ORF02334         SAG2103         563         arginyl-tRNA synthetase           ORF02337         SAG2105         290         conserved hypothetical protein           ORF02338         SAG2106         314         conserved hypothetical protein		SAG2090	136	
ORF02321         SAG2092         132         conserved hypothetical protein           ORF02322         SAG2093         379         recA protein           ORF02323         SAG2094         competence/damage-inducible protein CinA FRAMESHIFT           ORF02325         SAG2095         183         DNA-3-methyladenine glycosylase I           ORF02327         SAG2096         196         Holliday junction DNA helicase RuvA           ORF02328         SAG2097         418         transporter, putative           ORF02329         SAG2098         659         DNA mismatch repair protein HexB           ORF02330         SAG2099         33         hypothetical protein           ORF02331         SAG2100         67         cold shock protein, CSD family           ORF02332         SAG2101         858         DNA mismatch repair protein HexA           ORF02333         SAG2102         145         arginine repressor ArgR, putative           ORF02334         SAG2103         563         arginyl-tRNA synthetase           ORF02337         SAG2105         290         conserved hypothetical protein           ORF02338         SAG2106         314         conserved hypothetical protein           ORF02339         SAG2107         583         aspartyl-tRNA synthetase </td <td>ORF02320</td> <td>SAG2091</td> <td>88</td> <td>conserved hypothetical protein</td>	ORF02320	SAG2091	88	conserved hypothetical protein
ORF02323         SAG2094         competence/damage-inducible protein CinA FRAMESHIFT           ORF02325         SAG2095         183         DNA-3-methyladenine glycosylase I           ORF02327         SAG2096         196         Holliday junction DNA helicase RuvA           ORF02328         SAG2097         418         transporter, putative           ORF02329         SAG2098         659         DNA mismatch repair protein HexB           ORF02330         SAG2099         33         hypothetical protein           ORF02331         SAG2100         67         cold shock protein, CSD family           ORF02332         SAG2101         858         DNA mismatch repair protein HexA           ORF02333         SAG2102         145         arginine repressor ArgR, putative           ORF02334         SAG2103         563         arginyl-tRNA synthetase           ORF02337         SAG2104         102         conserved hypothetical protein           ORF02338         SAG2105         290         conserved hypothetical protein           ORF02339         SAG2107         583         aspartyl-tRNA synthetase           ORF02340         SAG2108         426         histidyl-tRNA synthetase		SAG2092	132	
ORF02325 SAG2095 183 DNA-3-methyladenine glycosylase I ORF02327 SAG2096 196 Holliday junction DNA helicase RuvA ORF02328 SAG2097 418 transporter, putative ORF02329 SAG2098 659 DNA mismatch repair protein HexB ORF02330 SAG2099 33 hypothetical protein ORF02331 SAG2100 67 cold shock protein, CSD family ORF02332 SAG2101 858 DNA mismatch repair protein HexA ORF02333 SAG2102 145 arginine repressor ArgR, putative ORF02334 SAG2103 563 arginyl-tRNA synthetase ORF02335 SAG2104 102 conserved hypothetical protein ORF02337 SAG2105 290 conserved hypothetical protein ORF02338 SAG2106 314 conserved hypothetical protein ORF02339 SAG2107 583 aspartyl-tRNA synthetase ORF02340 SAG2108 426 histidyl-tRNA synthetase		SAG2093	379	
ORF02327 SAG2096 196 Holliday junction DNA helicase RuvA ORF02328 SAG2097 418 transporter, putative ORF02329 SAG2098 659 DNA mismatch repair protein HexB ORF02330 SAG2099 33 hypothetical protein ORF02331 SAG2100 67 cold shock protein, CSD family ORF02332 SAG2101 858 DNA mismatch repair protein HexA ORF02333 SAG2102 145 arginine repressor ArgR, putative ORF02334 SAG2103 563 arginyl-tRNA synthetase ORF02335 SAG2104 102 conserved hypothetical protein ORF02337 SAG2105 290 conserved hypothetical protein ORF02338 SAG2106 314 conserved hypothetical protein ORF02339 SAG2107 583 aspartyl-tRNA synthetase ORF02340 SAG2108 426 histidyl-tRNA synthetase				
ORF02327         SAG2096         196         Holliday junction DNA helicase RuvA           ORF02328         SAG2097         418         transporter, putative           ORF02329         SAG2098         659         DNA mismatch repair protein HexB           ORF02330         SAG2099         33         hypothetical protein           ORF02331         SAG2100         67         cold shock protein, CSD family           ORF02332         SAG2101         858         DNA mismatch repair protein HexA           ORF02333         SAG2102         145         arginine repressor ArgR, putative           ORF02334         SAG2103         563         arginyl-tRNA synthetase           ORF02335         SAG2104         102         conserved hypothetical protein           ORF02337         SAG2105         290         conserved hypothetical protein           ORF02338         SAG2106         314         conserved hypothetical protein           ORF02340         SAG2107         583         aspartyl-tRNA synthetase           ORF02340         SAG2108         426         histidyl-tRNA synthetase		SAG2095	183	<del></del>
ORF02328         SAG2097         418         transporter, putative           ORF02329         SAG2098         659         DNA mismatch repair protein HexB           ORF02330         SAG2099         33         hypothetical protein           ORF02331         SAG2100         67         cold shock protein, CSD family           ORF02332         SAG2101         858         DNA mismatch repair protein HexA           ORF02333         SAG2102         145         arginine repressor ArgR, putative           ORF02334         SAG2103         563         arginyl-tRNA synthetase           ORF02335         SAG2104         102         conserved hypothetical protein           ORF02337         SAG2105         290         conserved hypothetical protein           ORF02338         SAG2106         314         conserved hypothetical protein           ORF02340         SAG2108         426         histidyl-tRNA synthetase		SAG2096	196	
ORF02329         SAG2098         659         DNA mismatch repair protein HexB           ORF02330         SAG2099         33         hypothetical protein           ORF02331         SAG2100         67         cold shock protein, CSD family           ORF02332         SAG2101         858         DNA mismatch repair protein HexA           ORF02333         SAG2102         145         arginine repressor ArgR, putative           ORF02334         SAG2103         563         arginyl-tRNA synthetase           ORF02335         SAG2104         102         conserved hypothetical protein           ORF02337         SAG2105         290         conserved hypothetical protein           ORF02338         SAG2106         314         conserved hypothetical protein           ORF02340         SAG2107         583         aspartyl-tRNA synthetase           ORF02341         SAG2108         426         histidyl-tRNA synthetase		SAG2097	418	
ORF02330         SAG2099         33         hypothetical protein           ORF02331         SAG2100         67         cold shock protein, CSD family           ORF02332         SAG2101         858         DNA mismatch repair protein HexA           ORF02333         SAG2102         145         arginine repressor ArgR, putative           ORF02334         SAG2103         563         arginyl-tRNA synthetase           ORF02335         SAG2104         102         conserved hypothetical protein           ORF02337         SAG2105         290         conserved hypothetical protein           ORF02338         SAG2106         314         conserved hypothetical protein           ORF02340         SAG2107         583         aspartyl-tRNA synthetase           ORF02341         SAG2108         426         histidyl-tRNA synthetase		SAG2098	659	
ORF02331         SAG2100         67         cold shock protein, CSD family           ORF02332         SAG2101         858         DNA mismatch repair protein HexA           ORF02333         SAG2102         145         arginine repressor ArgR, putative           ORF02334         SAG2103         563         arginyl-tRNA synthetase           ORF02335         SAG2104         102         conserved hypothetical protein           ORF02337         SAG2105         290         conserved hypothetical protein           ORF02338         SAG2106         314         conserved hypothetical protein           ORF02340         SAG2107         583         aspartyl-tRNA synthetase           ORF02341         SAG2108         426         histidyl-tRNA synthetase		SAG2099	33	
ORF02332         SAG2101         858         DNA mismatch repair protein HexA           ORF02333         SAG2102         145         arginine repressor ArgR, putative           ORF02334         SAG2103         563         arginyl-tRNA synthetase           ORF02335         SAG2104         102         conserved hypothetical protein           ORF02337         SAG2105         290         conserved hypothetical protein           ORF02338         SAG2106         314         conserved hypothetical protein           ORF02339         SAG2107         583         aspartyl-tRNA synthetase           ORF02340         SAG2108         426         histidyl-tRNA synthetase		SAG2100		
ORF02333         SAG2102         145         arginine repressor ArgR, putative           ORF02334         SAG2103         563         arginyl-tRNA synthetase           ORF02335         SAG2104         102         conserved hypothetical protein           ORF02337         SAG2105         290         conserved hypothetical protein           ORF02338         SAG2106         314         conserved hypothetical protein           ORF02339         SAG2107         583         aspartyl-tRNA synthetase           ORF02340         SAG2108         426         histidyl-tRNA synthetase		SAG2101	858	
ORF02334 SAG2103 563 arginyl-tRNA synthetase ORF02335 SAG2104 102 conserved hypothetical protein ORF02337 SAG2105 290 conserved hypothetical protein ORF02338 SAG2106 314 conserved hypothetical protein ORF02339 SAG2107 583 aspartyl-tRNA synthetase ORF02340 SAG2108 426 histidyl-tRNA synthetase		SAG2102	145	
ORF02335 SAG2104 102 conserved hypothetical protein ORF02337 SAG2105 290 conserved hypothetical protein ORF02338 SAG2106 314 conserved hypothetical protein ORF02339 SAG2107 583 aspartyl-tRNA synthetase ORF02340 SAG2108 426 histidyl-tRNA synthetase		SAG2103	563	
ORF02337 SAG2105 290 conserved hypothetical protein ORF02338 SAG2106 314 conserved hypothetical protein ORF02339 SAG2107 583 aspartyl-tRNA synthetase ORF02340 SAG2108 426 histidyl-tRNA synthetase		SAG2104	102	
ORF02338 SAG2106 314 conserved hypothetical protein ORF02339 SAG2107 583 aspartyl-tRNA synthetase ORF02340 SAG2108 426 histidyl-tRNA synthetase		SAG2105	290	
ORF02339 SAG2107 583 aspartyl-tRNA synthetase ORF02340 SAG2108 426 histidyl-tRNA synthetase		SAG2106	314	
ORF02340 SAG2108 426 histidyl-tRNA synthetase		SAG2107	583	
			426	
	ORF02341	SAG2109	60	

Table 32: C nversion fORFR fN s. with SAGR fNos.

ORF Ref No.	SAGxxxx Ref N .	aa	Ann tation
ORF02342	SAG2110	49	ribosomal protein L33
ORF02343	SAG2111	173	conserved hypothetical protein
ORF02344	SAG2112	494	site-specific recombinase, phage integrase family
ORF02345	SAG2113	82	conserved hypothetical protein
ORF02346	SAG2114	342	conserved hypothetical protein
ORF02347	SAG2115	143	hypothetical protein
ORF02349	SAG2116	151	conserved hypothetical protein
ORF02350	SAG2117	71	hypothetical protein
ORF02351	SAG2118	306	transcriptional regulator, Cro/Cl family
ORF02352	SAG2119	373	conserved domain protein
ORF02355	SAG2120	56	hypothetical protein
ORF02356	SAG2121	176	hypothetical protein
ORF02357	SAG2122	223	DNA-binding response regulator
ORF02358	SAG2123	454	sensor histidine kinase
ORF02359	SAG2124	517	membrane protein, putative
ORF02360	SAG2125	308	carbamate kinase
ORF02361	SAG2126	332	ornithine carbamoyltransferase
ORF02362	SAG2127	431	sensor histidine kinase
ORF02363	SAG2128	277	response regulator
ORF02364	SAG2129	240	amino acid ABC transporter, ATP-binding protein
ORF02365	SAG2130	504	amino acid ABC transporter, permease and amino acid binding protein
ORF02367	SAG2131	847	membrane protein, putative
ORF02368	SAG2132	247	conserved hypothetical protein
ORF02369	SAG2133	118	conserved hypothetical protein
ORF02370	SAG2134	772	membrane protein, putative
ORF02371	SAG2135	179	transcriptional regulator, TetR family, putative
ORF02372	SAG2136	98	conserved hypothetical protein
ORF02373	SAG2137	203	ribosomal protein S4
ORF02374	SAG2138	95	conserved hypothetical protein
ORF02375	SAG2139	451	replicative DNA helicase
ORF02376	SAG2140	150	ribosomal protein L9
ORF02377	SAG2141	660	DHH family protein
ORF02378	SAG2142	613	glucose inhibited division protein A
ORF02379	SAG2143	203	conserved hypothetical protein TIGR00427
ORF02380	SAG2144	373	tRNA (5-methylaminomethyl-2-thiouridylate)- methyltransferase
ORF02381	SAG2145	222	L-serine dehydratase, iron-sulfur-dependent, beta subunit
ORF02382	SAG2146	290	L-serine dehydratase, iron-sulfur-dependent, alpha subunit
ORF02383	SAG2147	234	conserved hypothetical protein
ORF02384	SAG2148	179	LysM domain protein
ORF02385	SAG2149	264	cobalt transport family protein
ORF02386	SAG2150	280	ABC transporter, ATP-binding protein
ORF02387	SAG2151	279	ABC transporter, ATP-binding protein, FRAMESHIFT
ORF02388	SAG2152	180	CDP-diacylglycerol-glycerol-3-phosphate 3-phosphatidyltransferase
ORF02389	SAG2153	427	peptidase, M16 family
ORF02390	SAG2154	414	conserved hypothetical protein
ORF02391	SAG2155	117	conserved hypothetical protein
	SAG2156	369	recF protein
OKF02392			
ORF02392 ORF02393	SAG2157	278	transporter, putative



## Tabl 32: Conv rsi n f ORF Ref Nos. with SAG Ref Nos.

ORF R f No.	SAGxxxx Ref N .	aa	Annotation
ORF02396	SAG2159	493	inosine-5'-monophosphate dehydrogenase
ORF02397	SAG2160	161	transcriptional regulator, ArgR family
ORF02398	SAG2161	226	transcriptional regulator, Crp/Fnr family
ORF02399	SAG2162	234	conserved hypothetical protein
ORF02400	SAG2163	410	arginine deiminase
ORF02401	SAG2164	136	acetyltransferase, GNAT family
ORF02402	SAG2165	337	ornithine carbamoyltransferase
ORF02403	SAG2166	475	arginine/ornithine antiporter
ORF02404	SAG2167	318	carbamate kinase
ORF02405	SAG2168	341	tryptophanyl-tRNA synthetase
ORF02406	SAG2169	230	conserved hypothetical protein
ORF02407	SAG2170	290	conserved hypothetical protein
ORF02408	SAG2171	539	ABC transporter, ATP-binding protein
ORF02409	SAG2172	859	ABC transporter, permease protein, putative
ORF02410	SAG2173	159	conserved hypothetical protein TIGR00246
ORF02411	SAG2174	409	serine protease
ORF02412	SAG2175	257	partitioning protein, ParB family
ORF02413	SAG0001	453	
ORF02415	SAG0001 SAG0002	378	chromosomal replication initiator protein DnaA  DNA polymerase III, beta subunit
ORF02416	SAG0002	293	
514 02410	3AG0003	293	diacylglycerol kinase catalytic domain protein, putative
ORF02417	SAG0004	65	conserved hypothetical protein
ORF02418	SAG0005	67	hypothetical protein
ORF02419	SAG0006	371	conserved hypothetical GTP-binding protein
ORF02420	SAG0007	191	peptidyl-tRNA hydrolase
ORF02421	SAG0008	1165	transcription-repair coupling factor
ORF02422	SAG0009	31	hypothetical protein
ORF02423	SAG0010	90	S4 domain protein
ORF02424	SAG0011	123	cell division protein DivIC, putative
ORF02425	SAG0012	44	conserved hypothetical protein
ORF02426	SAG0013	428	conserved hypothetical protein
ORF02427	SAG0014	424	MesJ/Ycf62 family protein
ORF02428	SAG0015	180	hypoxanthine-guanine phosphoribosyltransferase
	0.1000.0	.00	hypoxanamie-guarinie priosprioribosymansierase
ORF02429	SAG0016	658	cell division protein FtsH
ORF03000	SAG0157		Dnase-related protein, DEGENERATE
ORF03001	SAG0579	142	conserved hypothetical protein
ORF03002	SAG0580	111	conserved hypothetical protein, truncation
ORF03003	SAG0652		Tn5252, Orf 28 protein, degenerate
ORF03004	SAG0655	57	conserved hypothetical protein
ORF03005	SAG0662	101	cylX protein
ORF03006	SAG0917	83	Tn916, hypothetical protein
ORF03007	SAG0920	23	Tn916, hypothetical protein
ORF03008	SAG0922	61	Tn916, hypothetical protein
ORF03009	SAG0924	28	Tn916, tetM leader peptide
ORF03010	SAG0936	39	Tn916, hypothetical protein
ORF03011	SAG1484	48	ribosomal protein L33
ORF03012	SAG1857	119	HNH endonuclease family protein
ORF03013	SAG1883	128	conserved hypothetical protein
ORF03014	SAG2065	50	ribosomal protein L33
ORF03015	SAG2004	67	conjugal transfer protein, interruption-N
	UNUZUUT	<del></del>	Conjugar dansier protein, interruption-N
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		L	L

gi|13621326|gb|AAK33146.1| gi|13621327|gb|AAK33147.1| gi|13621328|gb|AAK33148.1| gi|13621329|gb|AAK33149.1| gi|13621330|gb|AAK33150.1| gi]13621331|gb]AAK33151.1| gi|13621332|gb|AAK33152.1| gi|13621333|gb|AAK33153.1| gi|13621334|gb|AAK33154.1| gi|13621335|gb|AAK33155.1| gi]13621337|gb]AAK33156.1| gi|13621340|gb|AAK33158.1| gi]13621341|gb|AAK33159.1[ gi]13621343|gb|AAK33160.1| gi|13621344|gb|AAK33161.1| gi|13621346|gb|AAK33163.1| gi|13621347|qb|AAK33164.1| gi|13621348|gb|AAK33165.1| gi|13621349|gb|AAK33166.1| gi|13621350|gb|AAK33167.1| gi|13621353|gb|AAK33169.1| gi|13621354|gb|AAK33170.1| gi|13621355|gb|AAK33171.1| gi|13621357|gb|AAK33173.1| gi|13621358|gb|AAK33174.1| gi|13621359|gb|AAK33175.1| gi|13621361|gb|AAK33176.1| gi|13621362|gb|AAK33177.1| gi|13621363|gb|AAK33178.1| gi|13621364|gb|AAK33179.1| gi|13621365|gb|AAK33180.1| gi|13621366|gb|AAK33181.1i gi|13621367|gb|AAK33182.1| gi|13621368|gb|AAK33183.1| gi|13621369|gb|AAK33184.1| gi|13621370|gb|AAK33185.1| gi|13621372|gb|AAK33186.1| gi|13621373|gb|AAK33187.1| gi|13621374|gb|AAK33188.1| gi|13621375|gb|AAK33189.1| gi|13621376|gb|AAK33190.1| gi|13621377|gb|AAK33191.1| gi|13621378|gb|AAK33192.1| gi|13621379|gb|AAK33193.1| gi|13621380|gb|AAK33194.1| gi|13621382|gb|AAK33196.1| gi|13621383|gb|AAK33197.1| gi]13621384[gb]AAK33198.1] gi|13621385|gb|AAK33199.1| gi|13621386|gb|AAK33200.1| gi|13621387|gb|AAK33201.1| gi|13621388|gb|AAK33202.1| gi|13621389|gb|AAK33203.1| gi|13621390|gb|AAK33204.1| gi|13621391|gb|AAK33205.1| gi|13621392|gb|AAK33206.1|

gi|13621393|gb|AAK33207.1| gi|13621394|gb|AAK33208.1| gi|13621397|gb|AAK33210.1| gi|13621398|gb|AAK33211.1| gi|13621399|gb|AAK33212.1| gi|13621401|gb|AAK33214.1| gi|13621403|gb|AAK33215.1| gi|13621404|gb|AAK33216.1| gi|13621405|gb|AAK33217.1| gi|13621407|qb|AAK33218.1| gi|13621408|gb|AAK33219.1| gi|13621409|gb|AAK33220.1| gi[13621413]gb[AAK33224.1[ gi|13621415|gb|AAK33226.1| gi|13621416|qb|AAK33227.11 gi|13621418|gb|AAK33229.1| gi|13621419|gb|AAK33230.1| gi|13621424|gb|AAK33234.1| gi|13621425|gb|AAK33235.1| gi|13621426|gb|AAK33236.1| gi|13621434|gb|AAK33243.1| gi[13621450]gb[AAK33258.1] gi|13621455|gb|AAK33262.1| gi|13621456|gb|AAK33263.1| gi]13621457[gb|AAK33264.1[ gi|13621467|gb|AAK33273.1| gi|13621468|gb|AAK33274.1| gi|13621469|gb|AAK33275.1| gi[13621470|gb|AAK33276.1] gi|13621471|gb|AAK33277.1| gi|13621472|gb|AAK33278.1| gi]13621473|gb|AAK33279.1| gi|13621476|gb|AAK33281.1| gi|13621477|gb|AAK33282.1| gi[13621478]gb[AAK33283.1] gi|13621480|gb|AAK33285.1| gi|13621481|gb|AAK33286.1| gi|13621491|gb|AAK33295.1| gi|13621494|gb|AAK33298.1| gi|13621496|gb|AAK33299.1| gi|13621501|gb|AAK33304.1| gi|13621502|gb|AAK33305.1| gi|13621505|gb|AAK33307.1| gi|13621506|gb|AAK33308.11 gi|13621507|gb|AAK33309.1| gi|13621510|gb|AAK33312.1| gi|13621511|gb|AAK33313.1| gi|13621513|gb|AAK33315.1| gi|13621516|gb|AAK33317.1| gi|13621518|gb|AAK33319.1| gi|13621521|gb|AAK33322.1| gi|13621522|gb|AAK33323.11 gi|13621523|gb|AAK33324.1| gi|13621524|gb|AAK33325.1| gi|13621525|gb|AAK33326.1| gi|13621527|gb|AAK33327.1|

gi|13621528|gb|AAK33328.1| gi|13621529|gb|AAK33329.1| gi|13621530|gb|AAK33330.1| gi|13621531|gb|AAK33331.1| gi|13621532|gb|AAK33332.11 gi|13621533|gb|AAK33333.1| gi|13621534|gb|AAK33334.1| gi|13621535|gb|AAK33335.1| gi|13621536|gb|AAK33336.1| gi|13621537|gb|AAK33337.1| gi|13621539|gb|AAK33338.1| gi|13621540|gb|AAK33339.1| gi|13621541|gb|AAK33340.1| gi|13621542|gb|AAK33341.1| gi|13621543|gb|AAK33342.1| gi|13621544|gb|AAK33343.1| gi|13621546|gb|AAK33345.1| gi|13621547|gb|AAK33346.1| gi|13621548|gb|AAK33347.1| gi|13621550|gb|AAK33348.1| gi[13621551]gb[AAK33349.1] gi|13621552|gb|AAK33350.1| gi|13621553|gb|AAK33351.1| gi|13621554|gb|AAK33352.11 gi|13621555|gb|AAK33353.1| gi|13621557|gb|AAK33355.1| gi]13621559|gb|AAK33356.1| gi|13621560|gb|AAK33357.1| gi|13621561|qb|AAK33358.11 gi|13621562|gb|AAK33359.1| gi]13621563|gb|AAK33360.1| gi|13621564|gb|AAK33361.1| gi|13621565|gb|AAK33362.11 gi|13621566|qb|AAK33363.11 gi|13621567|gb|AAK33364.1| gi|13621569|gb|AAK33365.1| gi|13621571|gb|AAK33367.1| gi|13621572|gb|AAK33368.1| gi|13621573|gb|AAK33369.1| gi|13621574|gb|AAK33370.1| gi|13621575|gb|AAK33371.1| gi|13621576|gb|AAK33372.1| gi|13621577|gb|AAK33373.11 gi]13621579]gb|AAK33374.1| gi|13621581|gb|AAK33376.1| gi|13621582|gb|AAK33377.1| gi|13621583|gb|AAK33378.1| gi|13621584|gb|AAK33379.1| gi|13621585|gb|AAK33380.1| gi|13621586|gb|AAK33381.1| gi[13621588|gb|AAK33383.1] gi|13621589|gb|AAK33384.1| gi|13621590|gb|AAK33385.1| gi|13621592|gb|AAK33386.1| gi|13621593|gb|AAK33387.1| gi|13621594|gb|AAK33388.1|



gi|13621595|gb|AAK33389.1| gi|13621596|gb|AAK33390.1| gi[13621597|gb|AAK33391.1| gi|13621598|gb|AAK33392.1| gi|13621599|gb|AAK33393.1| gi]13621600|gb|AAK33394.1| gi]13621602|gb|AAK33395.1| gi|13621603|gb|AAK33396.1| gi|13621604|gb|AAK33397.1| gi|13621605|gb|AAK33398.1| gi|13621606|gb|AAK33399.1| gi|13621607|gb|AAK33400.1| gi|13621608|gb|AAK33401.1| gi|13621609|gb|AAK33402.1| gi|13621611|gb|AAK33404.1| gi|13621614|gb|AAK33406.1| gi|13621615|gb|AAK33407.1| gi|13621616|gb|AAK33408.1| gi|13621617|gb|AAK33409.1| gi|13621618|gb|AAK33410.1| gi|13621619|gb|AAK33411.1| gi|13621620|gb|AAK33412.1| gi|13621621|gb|AAK33413.1| gi|13621622|gb|AAK33414.1| gi|13621623|gb|AAK33415.1| gi|13621624|gb|AAK33416.1| gi|13621625|gb|AAK33417.1| gi|13621627|gb|AAK33419.1| gi|13621629|gb|AAK33420.1| gi|13621630|gb|AAK33421.1| gi)13621631|gb|AAK33422.1| gi|13621633|gb|AAK33424.1| gi|13621634|gb|AAK33425.1| gi|13621636|gb|AAK33427.1| gi|13621637|gb|AAK33428.1| gi|13621638|gb|AAK33429.1| gi|13621640|gb|AAK33430.1| gi|13621642|gb|AAK33432.1| gi|13621644|gb|AAK33434.1| gi|13621645|gb|AAK33435.1| gi|13621647|gb|AAK33437.1| gi|13621648|gb|AAK33438.1| gi|13621650|gb|AAK33440.1| gi|13621651|gb|AAK33441.1| gi|13621652|gb|AAK33442.1| gi|13621657|gb|AAK33446.1| gi|13621658|gb|AAK33447.1| gi|13621660|gb|AAK33449.1| gi|13621670|gb|AAK33458.1| gi|13621671|gb|AAK33459.1| gi|13621672|gb|AAK33460.1| gi|13621675|gb|AAK33462.1| gi|13621676|gb|AAK33463.1| gi|13621678|gb|AAK33465.1| gi|13621680|gb|AAK33467.1| gi|13621681|gb|AAK33468.1|

gi|13621682|gb|AAK33469.1| gi|13621683|gb|AAK33470.1| gi|13621684|gb|AAK33471.1| gi]13621685|gb|AAK33472.1| gi)13621688|gb|AAK33474.1| gi]13621689|gb|AAK33475.1| gi|13621690|gb|AAK33476.1| gi|13621691|gb|AAK33477.1| gi|13621692|gb|AAK33478.1| gi|13621693|gb|AAK33479.1| gi|13621694|gb|AAK33480.1| gi|13621695|gb|AAK33481.1| gi|13621697|gb|AAK33483.1| gi|13621698|gb|AAK33484.1| gi|13621700|gb|AAK33485.1| gi|13621701|gb|AAK33486.1| gij13621702|gb|AAK33487.1| gi|13621714|gb|AAK33498.1| gi|13621715|gb|AAK33499.1| gi]13621717|gb|AAK33501.1| gi|13621718|gb|AAK33502.1| gi|13621719|qb|AAK33503.11 gi|13621720|gb|AAK33504.1| gi|13621726|gb|AAK33509.1| gi|13621727|gb|AAK33510.1| gi|13621729|gb|AAK33512.1| gi|13621730|gb|AAK33513.1| gi]13621731[gb|AAK33514.1] gi[13621732|gb|AAK33515.1| gi|13621733|gb|AAK33516.1| gi|13621734|gb|AAK33517.1| gi|13621735|gb|AAK33518.1| gi[13621736]gb[AAK33519.1] gi|13621741|gb|AAK33523.1| gi|13621742|gb|AAK33524.1| gi[13621743]gb[AAK33525.1] gi|13621744|gb|AAK33526.1| gi|13621745|gb|AAK33527.1| gi|13621747|gb|AAK33528.1| gi|13621756|gb|AAK33537.1| gi|13621773|gb|AAK33552.1| gi|13621774|gb|AAK33553.1| gi|13621775|gb|AAK33554.1| gi|13621777|gb|AAK33556.1| gi|13621778|gb|AAK33557.1| gi|13621779|gb|AAK33558.1| gi|13621781|gb|AAK33559.1| gi|13621782|gb|AAK33560.1| gi|13621785|gb|AAK33563.1| gi|13621786|gb|AAK33564.1| gi|13621787|gb|AAK33565.1| gi|13621788|gb|AAK33566.1| gi|13621789|gb|AAK33567.1| gi|13621790|gb|AAK33568.1| gi|13621793|gb|AAK33571.1| gi|13621794|gb|AAK33572,1|

gi|13621796|gb|AAK33573.1| gi|13621797|gb|AAK33574.1| gi|13621799|gb|AAK33576.1| gi|13621800|gb|AAK33577.1| gi|13621802|gb|AAK33579.1| gi|13621806|gb|AAK33583.1| gi|13621808|gb|AAK33584.1| gi|13621809|gb|AAK33585.1| gi|13621810|gb|AAK33586.1| gi|13621811|gb|AAK33587.1| gi|13621812|gb|AAK33588.1| gi|13621813|gb|AAK33589.1| gi|13621814|gb|AAK33590.1| gi|13621817|gb|AAK33592.1| gi|13621818|gb|AAK33593.1| gi|13621819|gb|AAK33594.1| gi|13621820|qb|AAK33595.1| gi|13621821|gb|AAK33596.1| gi]13621822]gb[AAK33597.1] gi|13621823|gb|AAK33598.1| gi|13621824|gb|AAK33599.1| gi|13621825|qb|AAK33600.1| gi|13621826|gb|AAK33601.1| gi|13621828|gb|AAK33602.1| gi|13621829|gb|AAK33603.1| gi|13621830|gb|AAK33604.1| gi|13621831|gb|AAK33605.1| gi|13621834|gb|AAK33608.1| gi[13621835|gb|AAK33609.1] gi|13621836|gb|AAK33610.1| gi|13621837|gb|AAK33611.1| gi|13621839|gb|AAK33612.1| gi|13621840|gb|AAK33613.1| gi|13621841|gb|AAK33614.1| gi[13621842|gb]AAK33615.1] gi|13621843|gb|AAK33616.1| gi|13621844|gb|AAK33617.1| gi|13621898|qb|AAK33667.1| gi|13621901|gb|AAK33670.1| gi[13621902|gb|AAK33671.1] gi|13621904|gb|AAK33672.1| gi|13621907|gb|AAK33675.1| gi|13621908|gb|AAK33676.1| gi|13621909|gb|AAK33677.1| gi]13621910|gb|AAK33678.1| gi[13621912]qb[AAK33680.1] gi]13621924|gb|AAK33690.1| gi|13621929|gb|AAK33694.1| gi|13621930|gb|AAK33695.1| gi|13621931|gb|AAK33696.1| gi|13621933|gb|AAK33698.1| gi|13621934|gb|AAK33699.1| gi|13621935|gb|AAK33700.1| gi|13621936|gb|AAK33701.1| gi|13621937|gb|AAK33702.1| gi|13621938|gb|AAK33703.1|

gi|13621939|gb|AAK33704.1| gi|13621942|gb]AAK33706.1| gi|13621944|gb|AAK33708.1| gi|13621945|gb|AAK33709.1| gi|13621946|gb|AAK33710.1| gij13621950|gb|AAK33714.1| gi|13621953|gb|AAK33716.1| gi|13621954|gb|AAK33717.1| gi|13621955|gb|AAK33718.1| gi]13621956|gb|AAK33719.1| gi|13621957|gb|AAK33720.1| gi|13621958|gb|AAK33721.1| gi]13621959|gb|AAK33722.1| gi]13621961|gb|AAK33723.1| gij13621975|gb|AAK33736.1| gi|13621977|gb|AAK33738.1| gi]13621978|qb|AAK33739.1| gi|13621979|gb|AAK33740.1| gi[13621980|gb|AAK33741.1| gi[13621981|gb|AAK33742.1] gi|13621982|gb|AAK33743.1| gi|13621985|gb|AAK33745.1| gi[13621986|gb|AAK33746.1] gi|13621987|gb|AAK33747.1| gi|13621989|gb|AAK33749.1| gi|13621990|gb|AAK33750.1| gi|13621992|gb|AAK33752.1| gi|13621993|gb|AAK33753.1| gi|13621994|gb|AAK33754.1| gi|13621996|gb|AAK33755.1| gi|13621997|gb|AAK33756.1| gi|13621998|gb|AAK33757.1| gi|13621999|gb|AAK33758.1| gi|13622000|gb|AAK33759.1| gi|13622001|gb|AAK33760.1| gi|13622002|gb|AAK33761.1| gi|13622003|gb|AAK33762.11 gi|13622004|gb|AAK33763.1| gi|13622005|gb|AAK33764.1| gi|13622006|gb|AAK33765.1| gi]13622008|gb|AAK33766.1| gi|13622009|gb|AAK33767.1| gi|13622010|gb|AAK33768.1| gi|13622012|gb|AAK33770.1| gi|13622013|gb|AAK33771.1| gi|13622017|gb|AAK33774.1| gi|13622018|gb|AAK33775.1| gi|13622019|gb|AAK33776.1| gi|13622020|gb|AAK33777.1| gi|13622021|gb|AAK33778.1| gi|13622024|gb|AAK33781.1| gi|13622025|gb|AAK33782.1| gi|13622026|gb|AAK33783.1| gi|13622031|gb|AAK33787.1| gi|13622032|gb|AAK33788.1| gi|13622033|gb|AAK33789.1|

gi|13622034|qb|AAK33790.11 gi|13622035|gb|AAK33791.1| gi|13622039|gb|AAK33794.1| gi|13622041|gb|AAK33796.1| gi|13622042|gb|AAK33797.1| gi|13622043|gb|AAK33798.1| gi|13622044|gb|AAK33799.1| gi|13622045|gb|AAK33800.1| gi|13622046|gb|AAK33801.1| gij13622048|gb|AAK33802.1| gi|13622049|gb|AAK33803.1| gi|13622050|gb|AAK33804.1| gi]13622051[gb]AAK33805.1] gi]13622052|gb|AAK33806.1| gi|13622054|gb|AAK33808.1| gi|13622055|gb|AAK33809.1| gi|13622056|gb|AAK33810.1| gi|13622058|gb|AAK33812.1| gi|13622060|gb|AAK33813.1| gi|13622062|gb|AAK33815.1| gij13622064[gb]AAK33817.1[ gi|13622065|gb|AAK33818.1| gi|13622068|qb|AAK33821.1| gi|13622069|gb|AAK33822.1| gi|13622070|gb|AAK33823.1| gi]13622071|gb|AAK33824.1| gi|13622073|gb|AAK33825.1| gi|13622074|gb|AAK33826.1| gi|13622075|qb|AAK33827.11 gi|13622077|gb|AAK33829.1| gi|13622079|gb|AAK33831.1| gi|13622083|gb|AAK33834.1| gi|13622085|gb|AAK33836.1| gi|13622086|qb|AAK33837.1| gi|13622087|gb|AAK33838.11 gi|13622088|gb|AAK33839.1| gi|13622089|gb|AAK33840.1| gi|13622090|gb|AAK33841.1| gi|13622091|qb|AAK33842.1| gi|13622092|gb|AAK33843.1| gi|13622093|gb|AAK33844.1| gi]13622095|gb|AAK33845.1| gi]13622096|gb|AAK33846.1| gi|13622097|gb|AAK33847.1| gi|13622162|gb|AAK33908.1| gi|13622163|gb|AAK33909.1| gi|13622164|gb|AAK33910.1| gi|13622165|gb|AAK33911.1| gi|13622166|gb|AAK33912.1| gi|13622169|gb|AAK33914.1| gi|13622170|gb|AAK33915.1| gi|13622171|gb|AAK33916.1| gi|13622172|gb|AAK33917.1| gi]13622174|gb|AAK33919.1| gi|13622175|gb|AAK33920.1| gi|13622176|gb|AAK33921.1|



gi|13622177|gb|AAK33922.1| gi|13622179|gb|AAK33923.1| gi|13622180|gb|AAK33924.1| gi]13622181|gb|AAK33925.1| gi|13622182|gb|AAK33926.1| gi|13622183|gb|AAK33927.1| gi[13622184]gb|AAK33928.1] gi|13622185|gb|AAK33929.1| gi|13622186|gb|AAK33930.1| gi|13622189|gb|AAK33932.1| gi|13622190|qb|AAK33933.1| gi|13622191|gb|AAK33934.1| gi|13622192|gb|AAK33935.1| gi|13622198|gb|AAK33940.1| gi|13622200|gb|AAK33942.1| gi|13622201|qb|AAK33943.1| gi|13622204|gb|AAK33946.1| gi|13622205|gb|AAK33947.1| gi|13622207|gb|AAK33949.1| gi|13622208|gb|AAK33950.1| gi|13622211|gb|AAK33952.1| gi|13622213|gb|AAK33954.1| gi]13622214|gb|AAK33955.1| gi]13622215|gb|AAK33956.1| gi|13622216|gb|AAK33957.1| gi|13622217|qb|AAK33958.1| gi|13622218|gb|AAK33959.1| gi]13622219|gb|AAK33960.1| gi|13622222|gb|AAK33962.1| gi]13622223|gb|AAK33963.1| gi[13622224]qb[AAK33964.1] gi|13622233|gb|AAK33972.1| gi|13622235|gb|AAK33974.1| gi|13622236|gb|AAK33975.1| gi|13622237|gb|AAK33976.1| gi|13622239|gb|AAK33978.1| gi|13622240|gb|AAK33979.1| gi|13622241|gb|AAK33980.1| gi|13622242|gb|AAK33981.1| gi|13622243|gb|AAK33982.1| gi]13622244|gb|AAK33983.1| gi|13622250|gb|AAK33988.1| gi|13622252|gb|AAK33990.1| gi|13622253|gb|AAK33991.1| gi|13622255|gb|AAK33993.1| gi|13622256|gb|AAK33994.1| gi|13622257|gb|AAK33995.1| gi|13622259|gb|AAK33996.1| gi[13622260]gb]AAK33997.1] gi[13622261]gb[AAK33998.1] gi|13622262|gb|AAK33999.1| gi|13622263|qb|AAK34000.1| gi|13622264|qb|AAK34001.1| gi|13622265|gb|AAK34002.1| gi|13622266|gb|AAK34003.1| gi|13622268|gb|AAK34005.1|

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gi|13622369|gb|AAK34097.1| gi|13622370|gb|AAK34098.11 gi|13622371|gb|AAK34099.11 gi|13622372|gb|AAK34100.1| gi|13622373|qb|AAK34101.11 gi|13622374|qb|AAK34102.11 gi]13622375|gb|AAK34103.1| gi|13622376|gb|AAK34104.1| gi|13622377|gb|AAK34105.1| gi|13622378|gb|AAK34106.1| gi|13622380|gb|AAK34107.1| gi|13622383|gb|AAK34110.1| gi|13622384|gb|AAK34111.1| gi|13622387|gb|AAK34114.1| gi|13622389|gb|AAK34116.1| gi|13622394|gb|AAK34120.1| gi|13622395|gb|AAK34121.1| gi|13622396|gb|AAK34122.1| gi|13622398|gb|AAK34124.1| gi|13622399|gb|AAK34125.1| gi|13622400|gb|AAK34126.1| gi|13622401|gb|AAK34127.1| gi|13622403|gb|AAK34128.1| gi|13622405|gb|AAK34130.1| gi|13622406|gb|AAK34131.1| gi|13622407[gb|AAK34132.1] gi]13622408|gb|AAK34133.1| gi|13622415|gb|AAK34139.1| gi|13622416|gb|AAK34140.11 gi|13622417|gb|AAK34141.1| gi|13622419|gb|AAK34143.1| gi|13622420|gb|AAK34144.1| gi|13622424|gb|AAK34147.1| gi|13622425|gb|AAK34148.1| gi|13622431|gb|AAK34153.1| gi|13622432|gb|AAK34154.1| gi|13622433|gb|AAK34155.1| gi|13622434|gb|AAK34156.1| gi|13622435|gb|AAK34157.1| gi|13622436|gb|AAK34158.1| gi|13622437|gb|AAK34159.1| gi|13622444|gb|AAK34165.1| gi|13622447|gb|AAK34168.1| gi|13622450|gb|AAK34170.1| gi|13622451|gb|AAK34171.1| gi|13622455|gb|AAK34175.1| gi|13622457|gb|AAK34177.1| gi|13622458|gb|AAK34178.1| gi|13622460|gb|AAK34179.1| gi|13622461|gb|AAK34180.1| gi|13622462|gb|AAK34181.1| gi|13622463|gb|AAK34182.1| gi|13622464|gb|AAK34183.1| gi|13622465|gb|AAK34184.1| gi|13622467|gb|AAK34186.1| gi|13622468|gb|AAK34187.1|

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gi|13622604|gb|AAK34311.1| gi|13622606|gb|AAK34313.1| gi|13622607|gb|AAK34314.1| gi|13622608|gb|AAK34315.1| gi|13622609|gb|AAK34316.1| gi|13622610]gb|AAK34317.1| gi|13622611|gb|AAK34318.1| gi[13622612]gb[AAK34319.1] gi]13622615|gb|AAK34321.1| gi|13622616|gb|AAK34322.1| gi|13622617|qb|AAK34323.1| gi|13622618|gb|AAK34324.1| gi]13622621|gb|AAK34327.1| gi|13622622]gb[AAK34328.1] gi|13622623|gb|AAK34329.1| gi]13622624|qb|AAK34330,1| gi|13622625|gb|AAK34331.1| gi|13622626|gb|AAK34332.1| gi|13622628|gb|AAK34333.1| gi|13622629|gb|AAK34334.1| gi|13622630|gb|AAK34335.1| gi|13622631|gb|AAK34336.1| gi|13622632|gb|AAK34337.1| gi|13622634|gb|AAK34339.1| gi|13622636|gb|AAK34341.1| gi]13622640|gb|AAK34344.1| gil13622641|gb|AAK34345.1| gi|13622652|gb|AAK34355.1| gi|13622653|gb|AAK34356.1| gi|13622654|gb|AAK34357.1| gi|13622656|qb|AAK34359.1| gi|13622660|gb|AAK34363.1| gi|13622665|gb|AAK34367.1| gi|13622668|gb|AAK34370.1| gi|13622675|gb|AAK34376.1| gi|13622676|gb|AAK34377.1| gil13622683|gb|AAK34383.1| gi|13622684]gb|AAK34384.1] gi|13622685|gb|AAK34385.1| gi|13622688|gb|AAK34387.1| gi|13622689|gb|AAK34388.1| gi|13622690|gb|AAK34389.1| gi|13622691|gb[AAK34390.1| gi|13622692|gb|AAK34391.1| gi|13622693|gb|AAK34392.1| gi|13622694|gb|AAK34393.1| gi|13622695|gb|AAK34394.1| gi|13622696|gb|AAK34395.1| gi|13622698|gb|AAK34396.1| gi|13622699|gb|AAK34397.1| gi|13622700|gb|AAK34398.1| gi|13622701|gb|AAK34399.1| gi|13622702|gb|AAK34400.1| gi|13622703|gb|AAK34401.1| gi|13622704|gb|AAK34402.1| gi|13622705|gb|AAK34403.1|



gi|13622711|gb|AAK34408.1| gi|13622713|gb|AAK34410.1| gi|13622714|gb|AAK34411.1| gi|13622715|gb|AAK34412.1| gi[13622718|gb|AAK34414.1| gi|13622719|gb|AAK34415.1| gi|13622720|gb|AAK34416.1| gi|13622721|gb|AAK34417.1| gi|13622722|gb|AAK34418.1| gi|13622723|gb|AAK34419.1| gi|13622727|gb|AAK34422.1| gi|13622728|gb|AAK34423.1| gi|13622729|gb|AAK34424.1| gi|13622730|gb|AAK34425.1| gi|13622731|gb|AAK34426.1| gi|13622733|gb|AAK34428.1| gi|13622734|gb|AAK34429.1| gi|13622735|gb|AAK34430.1| gi|13622736|gb|AAK34431.1| gi]13622737[gb]AAK34432.1] gi|13622740|gb|AAK34434.1| gi|13622741|gb|AAK34435.1| gi|13622742|gb|AAK34436.1| gi[13622744|gb|AAK34438.1| gi|13622745|gb|AAK34439.1| gi|13622746|gb|AAK34440.1| gi|13622749|gb|AAK34442.1| gi|13622750|gb|AAK34443.1| gi|13622751|gb|AAK34444.1| gi[13622752]gb]AAK34445.1| gi|13622753|gb|AAK34446.1| gi|13622754|gb|AAK34447.1| gi|13622760|gb|AAK34452.1| gi|13622762|gb|AAK34454.1| gi|13622763|gb|AAK34455.1| gi|13622764|gb|AAK34456.1| gi|13622765|gb|AAK34457.1| gi|13622766|gb|AAK34458.1| gi|13622767|gb|AAK34459.1| gi|13622768|gb|AAK34460.1| gi|13622770|gb|AAK34462.1| gi|13622771|gb|AAK34463.1| gi|13622774|gb|AAK34465.1| gi|13622775|gb|AAK34466.1| gi|13622776|gb|AAK34467.1| gi|13622777|gb|AAK34468.1| gi|13622778|gb|AAK34469.1| gi|13622779|gb|AAK34470.1| gi|13622780|gb|AAK34471.1| gi|13622781|gb|AAK34472.1| gi|13622782|gb|AAK34473.1| gi|13622783|qb|AAK34474.1| gi|13622785|gb|AAK34475.1| gi|13622787|gb|AAK34477.1| gi|13622789|gb|AAK34479.1| gi|13622790|gb|AAK34480.1|

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gi|13622791|gb|AAK34481.1| gi|13622792|gb|AAK34482.1| gi|13622793|gb|AAK34483.1| gi|13622794|gb|AAK34484.1| gi]13622795]gb[AAK34485.1] gij13622796|gb|AAK34486.1| gi|13622798|gb|AAK34487.1| gi|13622799|gb|AAK34488.1| gi|13622800|gb|AAK34489.1| gi|13622801|gb|AAK34490.1| gi|13622802|qb|AAK34491.1| gi|13622803|gb|AAK34492.1| gi]13622804|gb|AAK34493.1| gi|13622805|gb|AAK34494.1| gi|13622806|qb|AAK34495.1| gi[13622807]gb|AAK34496.11 gi|13622808|gb|AAK34497.1| gi|13622809|gb|AAK34498.1| gi]13622810|gb|AAK34499.1| gi|13622812|qb|AAK34500.1| gi|13622813|gb|AAK34501.1| gi|13622814|gb|AAK34502.1| gi|13622815|gb|AAK34503.1| gi|13622818|gb|AAK34506.1| gi|13622821|qb|AAK34509.1| gi|13622822|gb|AAK34510.1| gi|13622823|gb|AAK34511.1| gi|13622825|gb|AAK34512.1| gi|13622826|gb|AAK34513.1| gi|13622827|gb|AAK34514.1| gi|13622828|gb|AAK34515.1| gi]13622829|gb|AAK34516.1| gi]13622830|gb|AAK34517.1| gi|13622833|gb|AAK34520.1| gi|13622838|gb|AAK34524.1| gi|13622839|gb|AAK34525.1| gi|13622840|gb|AAK34526.1| gi|13622841|gb|AAK34527.1| gi|13622847|gb|AAK34532.1| gi|13622848|gb|AAK34533.1| gi|13622849|gb|AAK34534.1| gi|13622853|gb|AAK34537.1| gi|13622854|gb|AAK34538.1| gi[13622856]gb]AAK34540.1] gi|13622857|gb|AAK34541.1| gi|13622858|gb|AAK34542.1| gi|13622860|gb|AAK34543.1| gi|13622861|gb|AAK34544.1| gi|13622862|gb|AAK34545.1| gi|13622863|gb|AAK34546.1| gi|13622864|gb|AAK34547.1| gi|13622865|gb|AAK34548.1| gi|13622866|gb|AAK34549.1| gi|13622867|gb|AAK34550.1| gi|13622868|gb|AAK34551.1| gi|13622869|gb|AAK34552.1|

gi|13622870|gb|AAK34553.11 gi|13622873|gb|AAK34555.11 gi|13622875|gb|AAK34557.1| gi]13622876|gb|AAK34558.1| gi|13622877|gb|AAK34559.1| gi|13622878|gb|AAK34560.1| gi|13622879|gb|AAK34561.1| gi|13622880|gb|AAK34562.1| gi|13622881|gb|AAK34563.1| gi|13622882|gb|AAK34564.1| gi|13622885|gb|AAK34566.1| gi|13622886|gb|AAK34567.1| gi|13622887|gb|AAK34568.1| gi|13622888|gb|AAK34569.1| gi|13622890|gb|AAK34571.1| gi|13622893|gb|AAK34574.1| gi|13622896|gb|AAK34576.1| gi|13622898|gb|AAK34578.1| gi]13622899[gb]AAK34579.1] gi|13622900|gb|AAK34580.1| gi|13622901|gb|AAK34581.1| gi|13622903|gb|AAK34583.1| gi[13622905|qb|AAK34585.11 gi|13622906|gb|AAK34586.1] gi|13622907|gb|AAK34587.1| gi|13622908|gb|AAK34588.1| gi|13622910|gb|AAK34589.1| gi|13622911|gb|AAK34590.1| gi|13622912|ab|AAK34591.1| gi|13622913|gb|AAK34592.1| gi[13622914]gb[AAK34593.1] gi|13622915|gb|AAK34594.1| gi|13622917|gb|AAK34596.1| gi|13622918|qb|AAK34597.11 gi|13622919|gb|AAK34598.1| gi|13622921|gb|AAK34599.1| gi|13622922|gb|AAK34600.1| gi|13622924|qb|AAK34602.1| gi|13622925|gb|AAK34603.1| gi|13622926|gb|AAK34604.1| gi|13622927|gb|AAK34605.1| gi|13622928|gb|AAK34606.1| gi|13622929|gb|AAK34607.1| gi|13622930|gb|AAK34608.1| gi]13622931|gb|AAK34609.1| gi]13622933|gb|AAK34610.1| gi|13622941|gb|AAK34617.1| gi|13622944|gb|AAK34620.1| gi[13622945|gb|AAK34621.1| gi|13622947|gb|AAK34623.1| gi|13622948|gb|AAK34624.1| gi|13622949|gb|AAK34625.1| gi[13622950|gb|AAK34626.1] gi|13622952|gb|AAK34627.1| gi|13622955[gb|AAK34630.1] gi|13622956|gb|AAK34631.1|



gi|13622959|gb|AAK34634.1| gi|13622961|gb|AAK34636.1| gi|13622963|gb|AAK34638.1| gi|13622964|gb|AAK34639.1| gi|13622967|gb|AAK34641.1| gi|13622969|gb|AAK34643.1| gi|13622971|gb|AAK34645.1| gi|13622973|gb|AAK34647.1| gi|13622974|gb|AAK34648.1| gi|13622977|gb|AAK34651.1| gi|13622981|qb|AAK34654.1| gi|13622982|gb|AAK34655.1| gi|13622983|gb|AAK34656.1| gi|13622984|gb|AAK34657.1| gil13622985|ab|AAK34658.1| gi|13622989|gb|AAK34661.1| gi|13622990|gb|AAK34662.1| gi|13622991|gb|AAK34663.1| gi|13622992|gb|AAK34664.1| gi]13622995|gb|AAK34666.1| gi|13622996|gb|AAK34667.1| gi|13622998|gb|AAK34669.1| gi|13622999|gb|AAK34670.1| gi]13623000|gb|AAK34671.1| gi[13623001]gb[AAK34672.1] gi|13623002|gb|AAK34673.1| gi|13623004|gb|AAK34674.1| gi|13623005|gb|AAK34675.1| gi|13623006|gb|AAK34676.1| gi|13623007|gb|AAK34677.1| gi|13623009|gb|AAK34679.1| gi|13623019|gb|AAK34688.1| gi|13623020|gb|AAK34689.1| gi|13623030|gb|AAK34698.1| gi|13623031|gb|AAK34699.1| gi]13623032|gb|AAK34700.1| gi|13623033|gb|AAK34701.1| gi|13623038|gb|AAK34705.1| gi|13623045|gb|AAK34712.1| gi|13623046|gb|AAK34713.1| gi|13623047|gb|AAK34714.1| gi|13623049|gb|AAK34715.1| gi|13623050|gb|AAK34716.1| gi|13623051|gb|AAK34717.1| gi|13623052|gb|AAK34718.1| gi|13623053|gb|AAK34719.1| gi[13623054|gb|AAK34720.1| gi|13623056|gb|AAK34722.1| gi|13623058|gb|AAK34724.1| gi|13623062|gb|AAK34727.1| gi|13623064|gb|AAK34729.1| gi|13623065|gb|AAK34730.1| gi|13623069|gb|AAK34733.1| gi|13623074|gb|AAK34738.1| gi|13623081|gb|AAK34744.1| gi|13623082|gb|AAK34745.1|

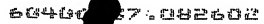


gi|13623083|gb|AAK34746.1| gi]13623085|gb|AAK34747.1| gi]13623086|gb|AAK34748.1| gi]13623088|gb|AAK34750.1| gi]13623089|gb|AAK34751.1| gi|13623090|gb|AAK34752.1| gi|13623091|gb|AAK34753.1| gi|13623093|gb|AAK34755.1| gi|13623095|gb|AAK34756.1| gi|13623096|gb|AAK34757.1| gi|13623098|qb|AAK34759.1| gi]13623099|gb|AAK34760.1| gi|13623100|gb|AAK34761.1| gi|13623102|gb|AAK34763.1| gi|13623103|gb|AAK34764.1| gi|13623105|qb|AAK34766.1| gi|13623107|gb|AAK34767.1| gi|13623128|gb|AAK34787.1| gi|13623129|gb|AAK34788.1| gi|13623131|gb|AAK34790.1| gi]13623132|qb|AAK34791.1| gi|13623133|gb|AAK34792.1| gi|13623134|gb|AAK34793.1| gi|13623136|gb|AAK34794.1| gi|13623138|gb|AAK34796.1| gi|13623139|gb|AAK34797.1| gi|13623150|gb|AAK34807.1| gi|13623151|gb|AAK34808.1| gi|13623152|gb|AAK34809.1| gi|13623154|gb|AAK34811.1| gi|13623155|gb|AAK34812,1| gi|13623156|gb|AAK34813.1| gi|13623157|gb|AAK34814.1| gi|13623159|gb|AAK34815.1| gi]13623161|gb|AAK34817.1| gi]13623162[gb|AAK34818.1] gi|13623163|gb|AAK34819.1| gi|13623165|gb|AAK34821.1| gi|13623166|gb|AAK34822.1| gi]13623167|gb|AAK34823.1| gi]13623168|gb|AAK34824.1| gi|13623170|gb|AAK34826.1| gi|13623171|gb|AAK34827.1| gi|13623175|gb|AAK34830.1| gi]13623176|gb|AAK34831.1| gi]13623177|gb|AAK34832.1| gi|13623179|gb|AAK34834.1| gi]13623180[gb[AAK34835.1] gi]13623182|gb|AAK34836.1| gi|13623183|gb|AAK34837.1| gi|13623184|gb|AAK34838.1| gi|13623185|gb|AAK34839.1| gi]13623186|gb|AAK34840.1| gi|13623187|gb|AAK34841.1|



# able 34: List of GAS ORF's which are shar d with GBS but which denot have homologues with Spn

gi|13621381|gb|AAK33195.1| gi|13621423|gb|AAK33233.1| gi|13621440|gb|AAK33249.1| gil13621443|gb|AAK33251.1| gi|13621453|gb|AAK33260.1| gi|13621454|gb|AAK33261.1| gi|13621479|gb|AAK33284.1| gi|13621482|gb|AAK33287.1| gi|13621492|gb|AAK33296.1| gi|13621493|gb|AAK33297.1| gi|13621497|gb|AAK33300.1| gi|13621498|gb|AAK33301.1| gi|13621512|gb|AAK33314.1| gi|13621514|gb|AAK33316.1| gi[13621556]gb]AAK33354.1] gi|13621570|gb|AAK33366.1| gi[13621587|gb|AAK33382.1| gi|13621610|gb|AAK33403.1| gi]13621613|gb|AAK33405.1| gi|13621626|gb|AAK33418.1| gi|13621632|gb|AAK33423.1| gi|13621635|gb|AAK33426.1| gi|13621643|gb|AAK33433.1| gi|13621655|gb|AAK33444.1| gi|13621656|gb|AAK33445.1| gi|13621659|gb|AAK33448.1| gi|13621673|gb|AAK33461.1| gi|13621686|gb|AAK33473.1| gi|13621696|gb|AAK33482.1| gi|13621703|gb|AAK33488.1| gi[13621712|gb|AAK33497.1] gi|13621728|gb|AAK33511.1| gi]13621738|gb|AAK33520.1| gi|13621739|gb|AAK33521.1| gi|13621740|gb|AAK33522.1| gi|13621772|gb|AAK33551.1| gi|13621776|gb|AAK33555.1| gi|13621791|gb|AAK33569.1| gi|13621798|gb|AAK33575.1| gi[13621801|gb|AAK33578.1| gi|13621803|gb|AAK33580.1| gi|13621804|gb|AAK33581.1| gi|13621832|gb|AAK33606.1| gi|13621833|gb|AAK33607.1| gi|13621896|gb|AAK33665.1| gi|13621897|gb|AAK33666.1| gi|13621906|gb|AAK33674.1| gi|13621911|gb|AAK33679.1| gi[13621949]gb]AAK33713.1] gi]13621951|gb|AAK33715.1| gi|13621962|gb|AAK33724.1| gi|13621963|gb|AAK33725.1| gi|13621964|gb|AAK33726.1| gi]13621971|gb|AAK33732.1| gi|13621976|gb|AAK33737.1| gi|13621983|gb|AAK33744.1|



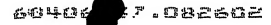
# ble 34: List of GAS ORF's which are shared with GBS but which denot have homel gues with Spn

gi|13621988|gb|AAK33748.1| gi|13622014|gb|AAK33772.1| gi|13622015|gb|AAK33773.1| gi|13622022|gb|AAK33779.1| gij13622023|gb|AAK33780.1| gi|13622028|gb|AAK33784.1| gi|13622029|gb|AAK33785.1| gi|13622037|gb|AAK33792.1| gi|13622038|gb|AAK33793.1| gi|13622040|gb|AAK33795.1| gi|13622057|gb|AAK33811.1| gi|13622061|gb|AAK33814.1| gi|13622063|gb|AAK33816.1| gi|13622066|gb|AAK33819.1| gi|13622067|gb|AAK33820.1| gi|13622076|gb|AAK33828.1| gi|13622078|gb|AAK33830.1| gi|13622084|gb|AAK33835.1| gi|13622098|gb|AAK33848.1| gi|13622099|gb|AAK33849.1| gi|13622100|gb|AAK33850.1| gi|13622104|gb|AAK33854.1| gi|13622110|gb|AAK33859.1| gi|13622116|gb|AAK33865.1| gi|13622124|gb|AAK33873.1| gi|13622159|gb|AAK33905.1| gi|13622193|gb|AAK33936.1| gi|13622194|gb|AAK33937.1| gi|13622195|gb|AAK33938.1| gi|13622196|gb|AAK33939.1| gi]13622202[gb]AAK33944.1] gil13622203|qb|AAK33945.11 gi|13622206|gb|AAK33948.1| gi|13622210|gb|AAK33951.1| gi|13622221|gb|AAK33961.1| gi]13622231|gb|AAK33971.1| gi|13622234|gb|AAK33973.1| gi|13622238|gb|AAK33977.1| gi]13622245|gb|AAK33984.1| gi|13622246|gb|AAK33985.1| gij13622248|gb|AAK33986.1| gi|13622249|gb|AAK33987.1| gi|13622251|gb|AAK33989.1| gi|13622254|gb|AAK33992.1| gi|13622267|gb|AAK34004.1| gi|13622291|gb|AAK34026.1| gi|13622302|gb|AAK34036.1| gi|13622303|gb|AAK34037.1| gi|13622304|gb|AAK34038.1| gi|13622327|gb|AAK34059.1| gi|13622344|gb|AAK34074.1| gi|13622345|gb|AAK34075.1| gi|13622346|gb|AAK34076.1| gi|13622347|gb|AAK34077.1| gi|13622348|gb|AAK34078.1| gi|13622349|gb|AAK34079.1|



# ble 34: List fGAS ORF's which are shared with GBS but which denot have homologues with Spn

gi|13622382|gb|AAK34109.1| gi|13622386|gb|AAK34113.1| gi|13622391|gb|AAK34118.1| gi|13622392|gb|AAK34119.1| gil13622397|gb|AAK34123.1| gi|13622404|gb|AAK34129.1| gi|13622412|gb|AAK34136.1| gi]13622413|gb|AAK34137.1| gi|13622414|gb|AAK34138.1| gi|13622418|gb|AAK34142.1| gi|13622430|gb|AAK34152.1| gi|13622446|gb|AAK34167.1| gi|13622449|gb|AAK34169.1| qi|13622453|qb|AAK34173.1| gi|13622470|gb|AAK34188.1| gi|13622487|gb|AAK34204.1| gi|13622490|gb|AAK34206.1| gi[13622502|gb|AAK34217.1| gi[13622503|gb|AAK34218.1| gi|13622514|gb|AAK34228.1| gi|13622528|gb|AAK34241.1| gi|13622540|gb|AAK34252.1| gi[13622541|gb|AAK34253.1| gi[13622544|gb|AAK34255.1| gi|13622545|gb|AAK34256.1| gi|13622546|gb|AAK34257.1| gi|13622547|gb|AAK34258.1| gi|13622548|gb|AAK34259.1| gi|13622550|gb|AAK34261.1| gi|13622551|gb|AAK34262.1| gi|13622552|gb|AAK34263.1| gi]13622556|gb|AAK34267.1| gi|13622557|gb|AAK34268.1| gi]13622558|gb|AAK34269.1| gi|13622559|gb|AAK34270.1| gi|13622563|gb|AAK34273.1| gi]13622571|gb|AAK34281.1| gi|13622576|gb|AAK34286.1| gi|13622581|gb|AAK34290.1| gi|13622582|gb|AAK34291.1| gi|13622586|gb|AAK34295.1| gi|13622589|gb|AAK34298.1| gi[13622605]gb]AAK34312.1] gi|13622633|gb|AAK34338.1| gi]13622635|gb]AAK34340.1| gi]13622637|gb|AAK34342.1| gij13622638|gb|AAK34343.1| gi|13622657|gb|AAK34360.1| gi|13622707|gb|AAK34404.1| gi|13622716|gb|AAK34413.1| gi|13622724|gb|AAK34420.1| gi[13622732|gb|AAK34427.1| gi|13622743|gb|AAK34437.1| gi|13622761|gb|AAK34453.1| gi|13622773|gb|AAK34464.1| gi|13622788|gb|AAK34478.1|





# 34: List f GAS ORF's which are shared with GBS but which do not hav homologues with Spn

gi|13622816|gb|AAK34504.1| gi|13622817|gb|AAK34505.1| gi|13622846|gb|AAK34531.1| gi|13622852|gb|AAK34536.1| gi|13622874|gb|AAK34556.1| gi|13622889|gb|AAK34570.1| gi]13622891[gb[AAK34572.1] gi|13622892|gb|AAK34573.1| gi[13622897]gb[AAK34577.1] gi|13622902|gb|AAK34582.1| gi|13622904|gb|AAK34584.1| gi[13622916]gb]AAK34595.1] gi|13622923|gb|AAK34601.1| gi|13622934|gb|AAK34611.1| gi[13622953]gb]AAK34628.1] gi]13622954|qb|AAK34629.1| gi[13622960]gb]AAK34635.1[ gi|13622968|gb|AAK34642.1| gi|13622980|gb|AAK34653.1| gi]13622987|gb|AAK34659.1| gi[13623012]gb]AAK34682.11 gi|13623013|gb|AAK34683.1| gi]13623014|gb|AAK34684.1| gi|13623015|gb|AAK34685.1| gi[13623016]gb]AAK34686.1] gij13623018jgbjAAK34687.11 gi]13623022|gb|AAK34691.1| gi]13623029|gb|AAK34697.1| gi|13623037|gb|AAK34704.1| gi|13623055|gb|AAK34721.1| gi]13623060|gb|AAK34725.1| gi|13623061|gb|AAK34726.1| gi|13623063|gb|AAK34728.1| gi[13623066]gb[AAK34731.1] gi|13623068|gb|AAK34732.1| gi|13623092|gb|AAK34754.1| gi|13623097|gb|AAK34758.1| gi|13623104|gb|AAK34765.1| gi]13623126]gb[AAK34785.1] gi|13623130|gb|AAK34789.1| gi|13623137|gb|AAK34795.1| gi|13623153|gb|AAK34810.1| gi|13623164|gb|AAK34820.1| gi|13623178|gb|AAK34833.1|



# Table 35: GAS ORF's which hav hom logu s with pneum coccus but which do not hav h m l gu s with GBS

gi|13621338|gb|AAK33157.1| gi|13621352|gb|AAK33168.1| gi|13621410|gb|AAK33221.1| gi|13621433|gb|AAK33242.1| gi|13621445|gb|AAK33253.1| gi|13621446|gb|AAK33254.1| gi]13621447|gb|AAK33255.1| gi|13621448|gb|AAK33256.1| gi|13621449|gb|AAK33257.1| gi|13621451|gb|AAK33259.1| gi|13621460|gb|AAK33267.1| gi|13621466|gb|AAK33272.1| gi|13621489|gb|AAK33293.1| gi|13621490|gb|AAK33294.1| gi|13621519|gb|AAK33320.1| gi|13621520|gb|AAK33321.1| gi|13621653|gb|AAK33443.1| gi|13621722|gb|AAK33506.1| gi[13621723]gb[AAK33507.1] gi[13621724|gb|AAK33508.1| gi|13621805|gb|AAK33582.1| gi|13621900|gb|AAK33669.1| gi|13622011|gb|AAK33769.1| gi[13622212]gb[AAK33953.1] gi[13622280]gb|AAK34016.11 gi|13622381|gb|AAK34108.1| gi[13622409]gb]AAK34134.1] gi|13622410|gb|AAK34135.1| gi|13622423|gb|AAK34146.1| gi]13622428|gb|AAK34151.1| gi|13622441|gb|AAK34162.1| gi|13622442|gb|AAK34163.1| gi|13622454|gb|AAK34174.1| gi|13622456|gb|AAK34176.1| gi|13622619|gb|AAK34325.1| gi|13622642|gb|AAK34346.1| gi|13622643|gb|AAK34347.1| gi|13622664|gb|AAK34366.1| gi|13622666|gb|AAK34368.1| gi|13622667|gb|AAK34369.1| gi|13622671|gb|AAK34372.1| gi|13622672|gb|AAK34373.1| gi|13622673|gb|AAK34374.1| gi|13622674|gb|AAK34375.1| gi|13622679|gb|AAK34380.1| gi|13622680|gb|AAK34381.1| gi|13622682|gb|AAK34382.1| gi|13622755|gb|AAK34448.1| gi|13622758|gb|AAK34450.1| gi|13622759|gb|AAK34451.1| gi|13622835|gb|AAK34521.1| gi|13622837|gb|AAK34523.1| gi|13622937|gb|AAK34614.1| gi|13622942|gb|AAK34618.1| gi]13622946|gb|AAK34622.1| gi|13622978|gb|AAK34652.1|



#### Table 35: GAS ORF's which hav homologues with pneum coccus but which do not hav h mologu s with GBS

gi|13623027|gb|AAK34695.1|

gi]13623087|gb|AAK34749.1|

gi|13623101|gb|AAK34762.1|

gi|13623144|gb|AAK34802.1|

gi|13623146|gb|AAK34804.1|

gi|13623147|gb|AAK34805.1|

Table 36: Spn ORF's which share homologues with GBS and GAS.

SP0001 SP0002 SP0003 SP0004 SP0005 SP0006 SP0007 SP0008 SP0010 SP0011 SP0013

SP0013 SP0014 SP0019 SP0021 SP0024

SP0024 SP0027 SP0032 SP0033 SP0034

SP0035 SP0036 SP0037 SP0042

SP0042 SP0044 SP0045

SP0046 SP0047

SP0048 SP0051 SP0053

SP0054 SP0056 SP0063

SP0073 SP0074

SP0078 SP0079 SP0083

SP0084 SP0085

SP0095 SP0105

SP0106 SP0111

SP0112

SP0118

SP0120 SP0121

SP0122

SP0127

SP0128 SP0129

SP0148

SP0149

SP0151 SP0152



Tabl 36: Spn ORF's which share h m logu s with GBS and GAS.

SP0158 SP0173 SP0179 SP0180 SP0184 SP0185 SP0186 SP0187 SP0189 SP0192 SP0194 SP0197 SP0199 SP0202 SP0204 SP0205 SP0208 SP0209 SP0210 SP0211 SP0212 SP0213 SP0214 SP0215 SP0216 SP0217 SP0218 SP0219 SP0220 SP0221 SP0222 SP0224 SP0225 SP0226 SP0227 SP0228 SP0229 SP0230 SP0231 SP0232 SP0233 SP0234 SP0235 SP0236 SP0240 SP0242 SP0243 SP0245 SP0246 SP0247 SP0248 SP0249 SP0250 SP0251

SP0252 SP0253



Tabl 36: Spn ORF's which shar hom logues with GBS and GAS.

SP0254 SP0259 SP0261 SP0262 SP0263 SP0264 SP0265 SP0266 SP0268 SP0271 SP0272 SP0273 SP0274 SP0280 SP0281 SP0282 SP0283 SP0284 SP0285 SP0286 SP0287 SP0289 SP0290 SP0291 SP0292 SP0294 SP0295 SP0303 SP0310 SP0314 SP0317 SP0318 SP0319 SP0320 SP0321 SP0322 SP0323 SP0324 SP0325 SP0327 SP0330 SP0334 SP0336 SP0337 SP0338 SP0340 SP0342 SP0369 SP0370 SP0371 SP0373 SP0374 SP0381 SP0382 SP0383 SP0384

3



SP0385

SP0386

SP0387

SP0400

SP0401

SP0402

SP0403

SP0404

SP0405

SP0406

SP0408

SP0410

SP0411

SP0412

SP0415

SP0416

SP0417

SP0418

SP0419

SP0420

SP0421

SP0422

SP0423

SP0424

SP0425

SP0426

SP0427

SP0433

SP0434

SP0435

SP0436

SP0437

SP0438

SP0439

SP0441 SP0442

SP0443

SP0452

SP0453

SP0454

SP0457

SP0458

SP0459

SP0461

SP0466

SP0467

SP0474

SP0477

SP0478

SP0483

SP0486

SP0488

SP0489

SP0493

SP0494

Tabl 36: Sph ORF's which shar h mologues with GBS and GAS.

SP0500 SP0501 SP0502 SP0515 SP0516 SP0517 SP0519 SP0521 SP0522 SP0523 SP0526 SP0549 SP0550 SP0552 SP0553 SP0554 SP0555 SP0556 SP0557 SP0563 SP0567 SP0568 SP0576 SP0577 SP0578 SP0579 SP0581 SP0588 SP0589 SP0591 SP0592 SP0593 SP0603 SP0604 SP0605 SP0608 SP0610 SP0611 SP0613 SP0614 SP0615 SP0616 SP0618 SP0620 SP0622 SP0623 SP0624 SP0626 SP0630 SP0631 SP0636 SP0637 SP0638 SP0645 SP0646



Table 36: Spn ORF's which shar homologues with GBS and GAS.

SP0787

SP0788 SP0792 SP0793 SP0797 SP0798 SP0799 SP0801 SP0802 SP0803 SP0805 SP0806 SP0807 SP0816 SP0817 SP0820 SP0822 SP0823 SP0824 SP0825 SP0828 SP0829 SP0831 SP0835 SP0837 SP0838 SP0839 SP0841 SP0843 SP0844 SP0845 SP0846 SP0847 **SP0848** SP0851 SP0852 SP0855 SP0856 SP0862 SP0864 SP0865 SP0867 SP0868 SP0869 SP0870 SP0871 SP0872 SP0873 SP0875 SP0876 SP0877 SP0878 SP0880 SP0881 SP0893 SP0894



#### Tabl 36: Spn ORF's which share homologues with GBS and GAS.

SP0895 SP0896 SP0897 SP0904 SP0905 SP0908 SP0909 SP0912 SP0923 SP0927 SP0928 SP0929 SP0931 SP0932 SP0933 SP0935 SP0936 SP0937 SP0938 SP0943 SP0944 SP0945 SP0946 SP0947 SP0948 SP0954 SP0955 SP0959 SP0960 SP0961 SP0962 SP0964 SP0966 SP0967 SP0968 SP0969 SP0970 SP0971 SP0972 SP0974 SP0975 SP0976 SP0978 SP0979 SP0980 SP0981 SP0984 SP0985 SP0987 SP0988 SP0989 SP0991 SP0992 SP0993 SP1002

6040 87.002602

Table 36: Spn ORF's which share homologues with GBS and GAS.

SP1004 SP1008 SP1010 SP1012 SP1016 SP1017 SP1018 SP1020 SP1021 SP1022 SP1024 SP1025 SP1026 SP1029 SP1033 SP1034 SP1035 SP1045 SP1056 SP1067 SP1068 SP1069 SP1070 SP1071 SP1072 SP1073 SP1074 SP1076 SP1079 SP1081 SP1082 SP1083 SP1084 SP1087 SP1088 SP1089 SP1090 SP1093 SP1094 SP1095 SP1096 SP1097 SP1098 SP1099 SP1100 SP1102 SP1105 SP1106 SP1107 SP1110 SP1111 SP1112 SP1113 SP1114 SP1115 SP1116

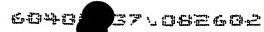




SP1117 **SP1118** SP1119 SP1128 SP1151 SP1152 SP1155 SP1156 SP1157 SP1159 SP1160 SP1161 SP1162 SP1163 SP1164 SP1167 SP1168 SP1169 SP1174 SP1175 SP1176 SP1177 SP1178 SP1179 SP1180 SP1182 SP1184 SP1185 SP1187 SP1190 SP1191 SP1192 SP1193 SP1197 SP1200 SP1202 SP1204 SP1205 SP1207 SP1208 SP1212 SP1213 SP1218 SP1219 SP1220 SP1225 SP1226 SP1227 SP1228 SP1229 SP1230 SP1231 SP1232 **SP1233** 

10

SP1238 SP1241



Tabl 36: Spn ORF's which share homol gues with GBS and GAS.

SP1242 SP1244 SP1245 SP1246 SP1247 **SP1248** SP1249 SP1260 SP1263 SP1266 SP1275 SP1276 SP1277 **SP1278** SP1279 SP1280 SP1283 SP1284 SP1285 SP1286 SP1287 SP1288 SP1289 SP1290 SP1291 SP1293 SP1297 SP1298 SP1299 SP1308 SP1316 SP1324 SP1329 SP1330 SP1331 SP1336 SP1341 SP1354 SP1355 SP1357 SP1358 SP1359 SP1362 SP1368 SP1370 SP1371 SP1372 **SP1374** SP1375 SP1376 SP1377 SP1378 SP1380 SP1381

SP1383 SP1386

Tabl 36: Sph-ORF's which share h mologu s with GBS and GAS.

SP1387 SP1388 SP1389 SP1390 SP1393 SP1394 SP1395 SP1396 SP1397 SP1398 SP1399 SP1400 SP1402 SP1403 SP1404 SP1405 SP1406 SP1407 SP1408 SP1409 SP1411 SP1412 SP1413 SP1414 SP1415 SP1416 SP1420 SP1421 SP1427 SP1428 SP1429 SP1434 SP1435 SP1445 SP1446 SP1448 SP1449 SP1450 SP1452 SP1453 SP1456 SP1457 SP1458 SP1460 SP1461 SP1462 SP1465 SP1466 SP1469 SP1470 SP1473 SP1474 SP1475 **SP1478** SP1479

Tabl 36: Spn ORF's which share homologu s with GBS and GAS.

SP1483 SP1485 SP1489 SP1491 SP1498 SP1500 SP1501 SP1502 SP1504 SP1505 SP1507 SP1508 SP1509 SP1510 SP1511 SP1512 SP1513 SP1517 SP1518 SP1519 SP1521 SP1522 SP1523 SP1529 SP1530 SP1534 SP1535 SP1536 SP1537 SP1538 SP1539 SP1540 SP1541 SP1542 SP1544 SP1547 SP1549 SP1551 SP1552 SP1553 SP1554 SP1557 SP1558 SP1559 SP1560 SP1561 SP1563 SP1564 SP1565 SP1566 SP1568 SP1569 SP1571 SP1574 SP1575



Tabl 36: Spn ORF's which share homologues with GBS and GAS.

SP1580 SP1583 SP1584 **SP1586** SP1587 **SP1588** SP1589 SP1590 SP1591 SP1597 SP1598 SP1599 SP1602 SP1603 SP1606 SP1608 SP1609 SP1610 SP1615 SP1616 SP1617 SP1624 SP1625 SP1626 SP1631 SP1633 SP1638 SP1644 SP1645 SP1646 SP1647 SP1648 SP1649 SP1650 SP1652 SP1653 SP1655 SP1659 SP1661 SP1662 SP1664 SP1665 SP1666 SP1667 SP1668 SP1670 SP1671 SP1672 SP1674 SP1675 SP1676 **SP1677** SP1681 SP1682 SP1683 SP1684



SP1685 SP1688 SP1689 SP1697 SP1698 SP1699 SP1702 SP1709 SP1711 SP1712 SP1713 SP1714 SP1717 SP1721 SP1722 SP1724 SP1725 SP1726 SP1727 SP1732 SP1733 SP1734 SP1735 SP1736 SP1737 **SP1738** SP1739 SP1742 SP1743 **SP1744** SP1746 SP1747 SP1748 SP1749 SP1750 SP1752 SP1759 SP1776 SP1780 SP1781 SP1782 SP1785 SP1790 SP1795 SP1799 SP1804 SP1816 SP1817 SP1825 SP1839 SP1840 SP1845 **SP1847** SP1848 SP1851 SP1855



Tabl 36: Spn ORF's which share h mologu s with GBS and GAS.

SP1857 SP1858 SP1860 SP1861 SP1865 SP1871 SP1873 SP1874 SP1875 SP1876 SP1877 SP1878 SP1879 SP1880 SP1881 SP1883 SP1884 **SP1887** SP1888 SP1889 SP1890 SP1895 SP1896 SP1900 SP1901 SP1902 SP1903 SP1906 SP1908 SP1909 SP1916 **SP1918** SP1922 SP1940 SP1942 SP1944 SP1953 SP1957 SP1960 SP1961 SP1963 SP1964 SP1966 SP1967 SP1968 SP1969 SP1970 SP1972 SP1973 SP1974 SP1975 SP1976 SP1979 SP1980 SP1981

Tabl 36: Spn ORF's which shar homologu s with GBS and GAS.

SP1983 SP1984 SP1985 SP1987 SP1989 SP1990 SP1991 SP1993 SP1994 SP1996 SP1997 SP1998 SP1999 SP2006 SP2007 SP2010 SP2011 SP2012 SP2020 SP2021 SP2022 SP2027 SP2028 SP2030 SP2031 SP2032 SP2033 SP2034 SP2035 SP2036 SP2037 **SP2038** SP2040 SP2041 SP2042 SP2044 SP2045 SP2048 SP2052 SP2053 SP2054 SP2055 SP2056 SP2057 SP2058 SP2063 SP2065 SP2069 SP2070 SP2072 SP2073 SP2075 SP2077 **SP2078** SP2082 SP2083



Tabl 36: Spn ORF's which shar homologu s with GBS and GAS.

SP2085 SP2086 SP2087 SP2088 SP2090 SP2091 SP2092 SP2094 SP2099 SP2100 SP2101 SP2106 SP2107 SP2108 SP2109 **SP2110** SP2112 SP2113 SP2114 SP2119 SP2121 SP2129 SP2131 SP2135 SP2142 SP2148 SP2150 SP2151 SP2152 SP2153 SP2156 SP2161 SP2162 SP2169 SP2170 SP2171 SP2172 SP2173 SP2174 SP2175 SP2176 SP2184 SP2185 SP2186 SP2187 SP2188 SP2189 SP2191 SP2192 SP2193 SP2194 SP2195 SP2202 SP2203 SP2204



#### Tabl 36: Spn ORF's which shar homol gu s with GBS and GAS.

SP2206 SP2207 SP2208 SP2209 SP2210 SP2214 SP2215 SP2216 SP2219 SP2220 SP2221 SP2222 SP2224 SP2225 SP2226 SP2227 SP2228 SP2229 SP2230 SP2231 SP2233 SP2234 SP2235 SP2238 SP2239 SP2240



# Tabl 37: Spn ORF's which share homologues with GBS but do n t shar hom I gues with GAS

SP0012 SP0020 SP0039 SP0050 SP0082 SP0107 SP0113 SP0119 SP0146 SP0150 SP0175 SP0176 SP0177 SP0178 SP0237 SP0255 SP0260 SP0267 SP0278 SP0288 SP0346 SP0347 SP0348 SP0349 SP0366 SP0376 SP0413 SP0445 SP0462 SP0463 SP0479 SP0480 SP0482 SP0484 SP0537 SP0538 SP0566 SP0580 SP0585 SP0599 SP0600 SP0601 SP0606 SP0607 SP0609 SP0617 SP0627 SP0655 SP0656 SP0710 SP0711 SP0717 SP0718 SP0720 SP0723



# Tabl 37: Spn ORF's which shar hom logu s with GBS but d not share h m l gues with GAS

SP0725 SP0730 SP0739 SP0749 SP0750 SP0751 SP0752 SP0753 SP0754 SP0769 SP0789 SP0791 SP0826 SP0900 SP0913 SP0914 SP0939 SP0941 SP0942 SP0953 SP0973 SP0977 SP1011 SP1013 SP1027 SP1054 SP1055 SP1080 SP1086 SP1121 SP1122 SP1123 SP1124 SP1126 SP1127 SP1137 SP1166 SP1173 SP1194 SP1195 SP1215 SP1240 SP1256 SP1261 SP1271 SP1272 SP1273 SP1274 SP1306 SP1310 SP1332 SP1333 SP1334 SP1346 SP1348



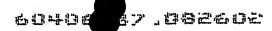
# Table 37: Spn ORF's which shar homologues with GBS but do not shar homol gues with GAS

SP1360 SP1361 SP1365 SP1382 SP1384 SP1392 SP1447 SP1451 SP1463 SP1464 SP1471 SP1472 SP1524 SP1527 SP1600 SP1605 SP1607 SP1632 SP1634 SP1651 SP1673 SP1680 SP1695 SP1700 SP1701 SP1720 SP1729 SP1740 SP1741 SP1745 SP1751 SP1757 SP1758 SP1761 SP1762 SP1763 SP1764 SP1765 SP1766 SP1767 SP1768 SP1770 SP1771 SP1772 SP1783 SP1802 SP1828 SP1856 SP1867 SP1869 SP1870 SP1872 SP1891 SP1907 SP1910



### Tabl 37: Spn ORF's which shar homol gues with GBS but do n t share h mologues with GAS

SP1927 SP1928 SP1943 SP1959 SP2001 SP2002 SP2009 SP2026 SP2029 SP2039 SP2061 SP2064 SP2066 SP2079 SP2084 SP2095 SP2096 SP2098 SP2103 SP2127 SP2128 SP2130 SP2134 SP2137 SP2138 SP2157 SP2196



#### Table 37: Spn ORF's which share homologues with GAS but do not share hom 1 gu s with GBS

SP0065 SP0075 SP0090 SP0091 SP0092 SP0099 SP0100 SP0153 SP0155 SP0156 SP0200 SP0306 SP0313 SP0341 SP0476 SP0496 SP0509 SP0527 SP0648 SP0658 SP0659 SP0661 SP0677 SP0715 SP0742 SP0743 SP0858 SP0859 SP0860 SP0910 SP0986 SP0994 SP0999 SP1000 SP1001 SP1023 SP1075 SP1129 SP1147 SP1171 SP1186 SP1315 SP1317 SP1319 SP1320 SP1321 SP1322 SP1438 SP1442 SP1525 SP1546

SP1570 SP1572 SP1578 SP1604 SP1715



#### Table 37: Spn ORF's which shar homologues with GAS but d not shar homol gues with GBS

SP1754 SP1797 SP1798 SP1800 SP1885 SP1919 SP1923 SP1941 SP1950 SP2016 SP2017 SP2051 SP2060 SP2111 SP2143 SP2144 SP2201 SP2236